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(71) Applicant: PROGENTIOR, INC. [US/US]; 4040 Campbell Avenue, Menlo Park, CA 94025 (US).

(72) Inventors: FEDER, John, N.; 1450 Chestnut Street, San Carlos. CA 94070 (US). KRONMAL, Gregory, S.; 277 Gateway Drive #131, Pacifica, CA 94044 (US). LAUER, Peter, M.; 128 Randall Street, San Francisco, CA 94131 (US). RUDDY, David, A.; 885 Greenwich Street, San Francisco, CA 94133 (US). THOMAS, Winston, J.; 40 White Plains Court, San Mateo, CA 94402 (US). TSUCHIHASHI, Zenta; 9 Light Way, Menlo Park, CA 94025 (US). WOLFF, Roger, K.; 41 Eugene Street, Mill Valley, CA 94941 (US).

(74) Agents: FITTS, Renee, A. et al.; Townsend and Townsend and Crew LLP, 8th floor, Two Embarcadero Center, San Francisco, CA 94111-3834 (US).

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(57) Abstract

Polymorphic sites in the region surrounding the HFE gene are provided. These polymorphisms are useful as surrogate markers in diagnostic assays for hemochromatosis. Additionally, a fine structure map of the 1 megabase region surrounding the HFE gene is provided, along with 235 kb of DNA sequence and 8 loci corresponding to candidate genes within the 1 megabase region, and in the purification of related proteins.

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# Polymorphisms and New Genes in the Region of the Human H mochromatosis Gene

# BACKGROUND OF THE INVENTION

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Hereditary hemochromatosis (HH) is an inherited disorder of iron metabolism wherein the body accumulates excess iron. In symptomatic individuals, this excess iron leads to deleterious effects by being deposited in a variety of organs leading to their failure, and resulting in cirrhosis, diabetes, sterility, and other serious illnesses. The gene which is defective in this disease was disclosed in copending U.S.S.N. 08/652,265.

Fine structure mapping of the region to which the gene responsible for HH, HFE (denoted HH or HFE in some publications), was mapped makes possible the identification of candidate sequences comprising the HFE gene, along with structural elements for regulation and expression and neighboring genes.

A variety of techniques is available for fine structure mapping, including direct cDNA selection, exon-trapping, and genomic sample sequencing. The direct selection approach (Lovett et al. Proc. Natl. Acad. Sci. U.S.A. 88:9628-9623 (1991)) involves the hybridization of cDNA fragments to genomic DNA. This technique is extremely sensitive and capable of isolating portions of rare transcripts. Exon-trapping (Church et al. Nature Genetics 6:98-105 (1994)) recovers spliced introns from in vivo expressed genomic DNA clones and produces candidate exons without requiring any prior knowledge of the target's gene expression. High-throughput genomic DNA sequencing with comparison of the sequence data to databases of expressed sequences has also been used, such as in the positional cloning of the Werner syndrome gene (Yu et al. Science 277:258-262 (1996)) and in cloning by homology of the second Alzheimer's disease gene on chromosome 1 (Levy-Lahad et al. Science 269:973-977 (1995)).

HH is typically inherited as a recessive trait; in the current state of knowledge, homozygotes carrying two defective copies of the gene are most frequently affected by the disease. In addition, heterozygotes for the HFE gene are more susceptible to sporadic porphyria cutanea tarda and potentially other disorders (Roberts et al., <u>Lancet</u> 349:321-323 (1997). It is estimated that approximately 10-15% of Caucasians carry one copy of the HFE gene mutation and that there are about one million homozygotes in the United States. HH, thus, represents one of the most common genetic disease mutations in Caucasian individuals. Although ultimately HH produces debilitating symptoms, the majority of homozygotes and heterozygotes have not been diagnosed.

The need for such diagnostics is documented, for example, in Barton, J.C. et al. Nature Medicine 2:394-395 (1996); Finch, C.A. West J Med 153:323-325 (1990); McCusick, V. Mendelian Inheritance in Man pp. 1882-1887, 11th ed., (Johns Hopkins University Press, Baltimore (1994)); Report of a Joint World Health Organization/Hemochromatosis Foundation/French Hemochromatosis Association Meeting on the Prevention and Control of Hemochromatosis (1993); Edwards, C.Q. et al. New Engl J Med 328:1616-1620 (1993); Bacon, B.R. New Engl J Med 326:126-

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127 (1992); Balan, V. et al. <u>Gastroenterology</u> 107:453-459 (1994); Phatak, P.D. et al. <u>Arch Int Med</u> 154:769-776 (1994).

A single mutation in the HFE gene, designated 24d1 in copending U.S.S.N. 08/630,912, gave rise to the majority of disease-causing chromosomes present in the population today. This is referred to herein as the "common" or "ancestral" or "common ancestral" mutation. These terms are used interchangeably. It appears that about 80% to 90% of all HH patients carry at least one copy of the common ancestral mutation which is closely linked to specific alleles of certain genetic markers close to this ancestral HFE gene defect. These markers are, as a first approximation, in the allelic form in which they were present at the time the ancestral HFE mutation occurred. See, for example, Simon, M. et al. Am J Hum Genet 41:89-105 (1987); Jazwinska, E.C. et al. Am J Hum Genet 53:242-257 (1993); Jazwinska, E.C. et al. Am J Hum Genet 56:428-433 (1995); Worwood, M. et al. Brit J Hematol 86:863-866 (1994); Summers, K.M. et al. Am J Hum Genet 45:41-48 (1989).

Several polymorphic markers in the HFE region have been described and shown to have alleles that are associated with HH disease. These markers include the published microsatellite markers D6S258, D6S306 (Gyapay, G. et al. Nature Genetics 7:246-339 (1994)), D6S265 (Worwood, M. et al. Brit J Hematol 86:833-846 (1994)), D6S105 (Jazwinska, E.C. et al. Am J Hum Genet 53:242-257 (1993); Jazwinska, E.C. et al. Am J Hum Genet 56:428-433 (1995)), D6S1001 (Stone, C. et al. Hum Molec Genet 3:2043-2046 (1994)), D6S1260 (Raha-Chowdhury et al. Hum Molec Genet 4:1869-1874 (1995)) as well as additional microsatellite and single-nucleotide-polymorphism markers disclosed in co-pending PCT application WO 96/06583, the disclosure of which is hereby incorporated by reference in its entirety. Additionally, copending U.S.S.N. 08/630,912 disclosed additional markers 24d2 and 24d7.

The symptoms of HH are often similar to those of other conditions, and the severe effects of the disease often do not appear immediately. Accordingly, it would be desirable to provide a method to identify persons who may be destined to become symptomatic in order to intervene in time to prevent excessive tissue damage associated with iron overload. One reason for the lack of early diagnosis is the inadequacy of presently available diagnostic methods to ascertain which individuals are at risk, especially while such individuals are presymptomatic.

Although blood iron parameters can be used as a screening tool, a confirmed diagnosis often employs liver biopsy which is undesirably invasive, costly, and carries a risk of mortality. Thus, there is a clear need for the development of an inexpensive and noninvasive diagnostic test for detection of homozygotes and heterozygotes in order to facilitate diagnosis in symptomatic individuals, provide presymptomatic detection to guide intervention in order to prevent organ damage, and for identification of heterozygote carriers.

Furthermore, a need exists for both methods for fine structure mapping and a fine structure map of the region of the chromosome to which the HH locus maps. This and other needs are addressed by the present invention.

## SUMMARY OF THE INVENTION

One aspect of the invention is an oligonucleotide comprising at least 8 to about 100 consecutive bases from the sequence of Figure 9, or the complement of the sequence, wherein the at least 8 to about 100 consecutive bases includes at least one polymorphic site of Table 1.

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Another aspect of the invention is an oligonucleotide pair selected from the sequence of Figure 9 or its complement for amplification of a polymorphic site of Table 1.

Another aspect of the invention is an isolated nucleic acid molecule comprising about 100 consecutive bases to about 235 kb substantially identical to the sequence of Figure 9, wherein the DNA molecule comprises at least one polymorphic site of Table 1.

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Another aspect of the invention is a method to determine the presence or absence of the common hereditary hemochromatosis (HFE) gene mutation in an individual comprising:

providing DNA or RNA from the individual; and

assessing the DNA or RNA for the presence or absence of a haplotype of

Table 1,

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wherein, as a result, the absence of a haplotype of Table 1 indicates the likely absence of the HFE gene mutation in the genome of the individual and the presence of the haplotype indicates the likely presence of the HFE gene mutation in the genome of the individual.

Another aspect of the invention is a method to determine the presence or absence of the common hereditary hemochromatosis (HFE) gene mutation in an individual comprising:

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providing DNA or RNA from the individual; and

assessing the DNA or RNA for the presence or absence of a genotype defined by a polymorphic allele of Table 1,

wherein, as a result, the absence of a genotype defined by a polymorphic allele of Table 1 indicates the likely absence of the HFE gene mutation in the genome of the individual and the presence of the genotype indicates the likely presence of the HFE gene mutation in the genome of the individual.

Another aspect of the invention is a culture of lymphoblastoid cells having the designation ATCC CRL-12371.

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One aspect of the invention is an isolated nucleic acid sequence comprising a nucleic acid sequence substantially identical to BTF1.

A further aspect of the invention is an isolated nucleic acid sequence comprising a nucleic acid sequence substantially identical to BTF2.

A further aspect of the invention is an isolated nucleic acid sequence comprising a nucleic acid sequence substantially identical to BTF3.

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A further aspect of the invention is an isolated nucleic acid sequence comprising a nucleic acid sequence substantially identical to BTF4.

A further aspect of the invention is an isolated nucleic acid sequence comprising a nucleic acid sequence substantially identical to BTF5.

A further aspect of the invention is an isolated nucleic acid sequence comprising a nucleic acid sequenc substantially identical to NPT3.

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A further aspect of the invention is an isolated nucleic acid's quence comprising a nucleic acid sequence substantially identical to NPT4.

A further aspect of the invention is an isolated nucleic acid sequence comprising a nucleic acid sequence substantially identical to RoRet.

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Additional aspects of the invention include nucleic acid sequences that are cDNAs, polypeptides encoded by the nucleic acids of the invention and antibodies specifically immunoreactive thereto, vectors comprising the nucleic acid sequences of the invention, and host cells stably transfected with the nucleic acids of the invention.

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A further aspect of the invention is an isolated nucleic acid sequence comprising at least 18 contiguous nucleotides substantially identical to at least 18 contiguous nucleotides of BTF1.

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A further aspect of the invention is an isolated nucleic acid sequence comprising at least 18 contiguous nucleotides substantially identical to at least 18 contiguous nucleotides of BTF2.

A further aspect of the invention is an isolated nucleic acid sequence comprising at least 18 contiguous nucleotides substantially identical to at least 18 contiguous nucleotides of BTF3.

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A further aspect of the invention is an isolated nucleic acid sequence comprising at least 18 contiguous nucleotides substantially identical to at least 18 contiguous nucleotides of BTF4.

A further aspect of the invention is an isolated nucleic acid sequence comprising at least 18 contiguous nucleotides substantially identical to at least 18 contiguous nucleotides of BTF5.

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A further aspect of the invention is an isolated nucleic acid sequence comprising at least 18 contiguous nucleotides substantially identical to at least 18 contiguous nucleotides of NPT3.

A further aspect of the invention is an isolated nucleic acid sequence comprising at least 18 contiguous nucleotides substantially identical to at least 18 contiguous nucleotides of NPT4.

A further aspect of the invention is an isolated nucleic acid sequence comprising at least 18 contiguous nucleotides substantially identical to at least 18 contiguous nucleotides of RoRet.

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# BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 depicts a combination genetic, physical and transcription map of the HFE gene region. The first line shows the relative positions of selected genetic markers that define the HFE region. The heavy bar below represents the YAC clone used in the direct selection experiment. The order and positions of the bacterial clones employed in the exon-trapping and sample sequencing is indicated under the YAC. The thin bar under the bacterial clones represents the approximate locations of a subset of the expressed sequence fragments mapped to the contig. The thicker bars show the location of the cDNAs cloned. Two regions are bracketed; the butyrophilin family of genes (BTF), and the region where complete genomic sequencing was carned out.

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Figure 2 is a schematic of the 250 kb of genomic sequence including the HFE gene. Both the structure of the overall cDNA (top) and that corresponding to the coding regions (bottom), as well as the direction of transcription are shown. The positions of the histone genes, the zinc  $\alpha$ -2 glycoprotein pseudogene, and the ESTs are also shown.

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Figure 3 depicts an alignment of the predicted amino acid sequence of the BTF proteins. Sequence were aligned in a pair-wise fashion using CLUSTAL W (Thompson et al. Nucl. Acids Res. 22:4573-4680) to deduce the most parsimonious arrangement. The ast risks under the

alignment represent amino acids conserved in all 6 proteins; the "dots" represent conserved amino acids substitutions. Box deare their gions within the proteins which correspond to three conserved metifs: 1) the B-G domain, 2) the transmembrane domain (TM), and 3) the B30-2 exon domain.

Figure 4, panel (A) depicts a Northern blot analysis of representative members of the two groups of BTF proteins, BTF1 and BTF5. BTF1 hybridized to all tissues on the blot as a major transcript at 2.9 kb and a minor one at 5.0 kb. BTF5 hybridized to several transcripts ranging between 4.0 and 3.1 kb and as a similar expression profile to BTF1. Autoradiography was for 24 hours. The β-actin hybridization demonstrated the variation in ploy (A)+ RNA between the lanes. Autoradiography was for 1 hour. In panel (B), RT-PCR analysis demonstrated that the expression of both genes was widespread. Included in the (+) lane are cDNA 21 and 44 as positive controls; the (-) lane represents the no-DNA control. Amplification using primers for the RFP gene (Isomura *et al.* <u>Nucleic Acid Res.</u> 20:5305-5310 (1992)) controlled for the integrity of the cDNA. All first strand cDNAs were checked for contaminating genomic DNA amplification by carrying out an identical experiment excluding the reverse transcriptase. In all cases, no amplification was obtained (data not shown).

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Figure 5(A) depicts an alignment of the predicted amino acid sequence of the RoRet gene to the 52 kD Ro/SSA auto-antigen protein. The asterisks under the alignment represent conserved amino acids; the "dots" represent conserved amino acids substitutions. The putative DNA binding cysteine-rich domain and the B30-2 exon domain are boxed. Figure 5(B) depicts an alignment of the predicted amino acid sequence of the two novel putative sodium phosphate transport proteins to that of the NPT1.

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Figure 6, panel (A) depicts a Northern blot analysis of the RoRet gene. The RoRet cDNA hybridized to 4 different transcripts, ranging from 7.1 kb to 2.2 kb. Autoradiography was performed for 4 days. The re-hybridization of the blot with a β-actin probe showed the variation in poly (A)+ RNA between the lanes. Autoradiography was for 1 hour. Panel (B) depicts RT-PCR analysis of the RoRet gene. Included in the (+) lane was a cDNA 27 positive control. Weak amplification of the correct size was observed in the small intestine, kidney and liver. The other tissues were negative as was the no DNA control lane (-). The RFP primers demonstrated the integrity of the cDNA. Panel (C) depicts Northern blot analysis of NPT3 and NPT4. NPT3 was expressed at high abundance in the heart and muscle as a single 7.2 kb transcript. Lesser amounts were found in the other tissues. The expression pattern of NPT4 was more restricted, being found only in the liver and kidney as a smear of transcripts ranging from 2.6 to 1.7 kb. Panel (D) depicts RT-PCR analysis of the NPT3 and NPT4 genes. Included in the (+) lane were the respective cDNA22E and 22B positive controls. The NPT3 gene was expressed as the proper size PCR fragment in kidney, liver, spleen and testis. A smaller fragment was detected in all tissues with the exception of the liver. The no DNA control lane (-) was negative. NPT4 was expressed as the proper size fragment in the small intestine, kidney, liver and testis. Larger and smaller size fragments were found in all other tissues with the exception of the brain. For both genes these different size fragments may indicate alternative splice events. The no DNA control lane (-) was negative. The RFP primers demonstrated the integrity of the cDNA.

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Figur 7 depicts the sequences of cDNA 21 (BTF1), cDNA 29 (BTF3), cDNA 23 (BTF4), cDNA 44 (BTF5), cDNA 32 (BTF2), cDNA 27 (RoRet), cDNA 22B (NPT3), cDNA22E (NPT4).

Figure 8 depicts the nucleotide sequence of approximately 235 kb in the HFE subregion from an unaffected individual.

Figure 9 depicts the nucleotide sequence of approximately 235 kb in the HFE subregion from an HH affected individual. Polymorphic sites in the HH affected individual determined by comparing a sequence of the corresponding region from an HH unaffected individual are listed and described in Table I.

#### **DETAILED DESCRIPTION**

### A. <u>Definitions</u>

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Abbreviations for the twenty naturally occurring amino acids follow conventional usage. In the polypeptide notation used herein, the left-hand direction is the amino terminal direction and the right-hand direction is the carboxyl-terminal direction, in accordance with standard usage and convention. Similarly, unless specified otherwise, the left hand end of single-stranded polynucleotide sequences is the 5' end; the left hand direction of double-stranded polynucleotide sequences is referred to as the 5' direction. The direction of 5' to 3' addition of nascent RNA transcripts is referred to as the transcription direction; sequence regions on the DNA strand having the same sequences: sequence regions on the DNA strand having the same sequence as the RNA and which are 5' to the 5' end of the RNA transcript are referred to as "upstream sequences"; sequence regions on the DNA strand having the same sequence as the RNA and which are 3' to the 3' end of the RNA transcript are referred to as "downstream sequences".

The term "nucleic acids", as used herein, refers to either DNA or RNA. "Nucleic acid sequence" or "polynucleotide sequence" refers to a single- or double-stranded polymer of deoxyribonucleotide or ribonucleotide bases read from the 5' to the 3' end. It includes both self-replicating plasmids, infectious polymers of DNA or RNA and nonfunctional DNA or RNA. The complement of any nucleic acid sequence of the invention is understood to be included in the definition of that sequence.

"Nucleic acid probes" may be DNA or RNA fragments. DNA fragments can be prepared, for example, by digesting plasmid DNA, or by use of PCR, or synthesized by either the phosphoramidite method described by Beaucage and Carruthers, <u>Tetrahedron Lett.</u> 22:1859-1862 (1981), or by the triester method according to Matteucci, et al., <u>J. Am. Chem. Soc.</u> 103:3185 (1981), both incorporated herein by reference. A double stranded fragment may then be obtained, if desired, by annealing the chemically synthesized single strands together under appropriate conditions or by synthesizing the complementary strand using DNA polymerase with an appropriate primer sequence. Where a specific sequence for a nucleic acid probe is given, it is understood that the complementary strand is also identified and included. The complementary strand will work equally well in situations where the target is a double-stranded nucleic acid.

The phrase "selectively hybridizing to" refers to a nucleic acid probe that hybridizes, duplexes or binds only to a particular target DNA or RNA sequence when the target sequences are present in a preparation of total cellular DNA or RNA. "Complementary" or "target" nucleic acid sequences refer to those nucleic acid sequences which selectively hybridize to a nucleic acid probe. Proper annealing conditions depend, for example, upon a probe's length, bas composition, and the number of mismatches and their position on the probe, and must often be determined empirically. For

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discussions of nucleic acid probe design and annealing conditions, see, for xample, Sambrook et al., Mol. cular Cloning: a Laboratory Manual (2nd ed.), Vols. 1-3, Cold Spring Harbor Laboratory, (1989) or Current Protocols in Molecular Biology, F. Ausubel et al., ed. Greene Publishing and Wiley-Interscience, New York (1987).

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The phrase "nucleic acid sequence encoding" refers to a nucleic acid which directs the expression of a specific protein or peptide. The nucleic acid sequences include both the DNA strand sequence that is transcribed into RNA and the RNA sequence that is translated into protein. The nucleic acid sequences include both the full length nucleic acid sequences as well as non-full length sequences derived from the full length protein. It being further understood that the sequence includes the degenerate codons of the native sequence or sequences which may be introduced to provide codon preference in a specific host cell.

The phrase "isolated" or "substantially pure" refers to nucleic acid preparations that lack at least one protein or nucleic acid normally associated with the nucleic acid in a host cell.

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The phrase "expression cassette", refers to nucleotide sequences which are capable of affecting expression of a structural gene in hosts compatible with such sequences. Such cassettes include at least promoters and optionally, transcription termination signals. Additional factors necessary or helpful in effecting expression may also be used as described herein.

The term "operably linked" as used herein refers to linkage of a promoter upstream from a DNA sequence such that the promoter mediates transcription of the DNA sequence.

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The term "vector", refers to viral expression systems, autonomous self-replicating circular DNA (plasmids), and includes both expression and nonexpression plasmids. Where a recombinant microorganism or cell culture is described as hosting an "expression vector," this includes both extrachromosomal circular DNA and DNA that has been incorporated into the host chromosome(s). Where a vector is being maintained by a host cell, the vector may either be stably replicated by the cells during mitosis as an autonomous structure, or is incorporated within the host's genome.

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The term "gene" as used herein is intended to refer to a nucleic acid sequence which encodes a polypeptide. This definition includes various sequence polymorphisms, mutations, and/or sequence variants wherein such alterations do not affect the function of the gene product. The term "gene" is intended to include not only coding sequences but also regulatory regions such as promoters, enhancers, and termination regions. The term further includes all introns and other DNA sequences spliced from the mRNA transcript, along with variants resulting from alternative splice sites.

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The term "plasmid" refers to an autonomous circular DNA molecule capable of replication in a cell, and includes both the expression and nonexpression types. Where a recombinant microorganism or cell culture is described as hosting an "expression plasmid", this includes both extrachromosomal circular DNA molecules and DNA that has been incorporated into the host chromosome(s). Where a plasmid is being maintained by a host cell, the plasmid is either being stably replicated by the cells during mitosis as an autonomous structure or is incorporated within the host's genome.

The phrase "recombinant protein" or "recombinantly produced protein" refers to a peptide or protein produced using non-native cells that do not have an endogenous copy of DNA able to express the protein. The cells produce the protein because they have been genetically altered by the introduction of the appropriate nucleic acid sequence. The recombinant protein will not be found in association with proteins and other subcellular components normally associated with the cells producing the protein. The terms "protein" and "polypeptide" are used interchangeably herein.

The following terms are used to describe the sequence relationships between two or more nucleic acids or polynucleotides: "reference sequence", "comparison window", "sequence identity", "percentage of sequence identity", and "substantial identity". A "reference sequence" is a defined sequence used as a basis for a sequence comparison; a reference sequence may be a subset of a larger sequence, for example, as a segment of a full-length cDNA or gene sequence given in a sequence listing, or may comprise a complete cDNA or gene sequence.

Optimal alignment of sequences for aligning a comparison window may, for example, be conducted by the local homology algorithm of Smith and Waterman Adv. Appl. Math. 2:482 (1981), by the homology alignment algorithm of Needleman and Wunsch J. Mol. Biol. 48:443 (1970), by the search for similarity method of Pearson and Lipman Proc. Natl. Acad. Sci. U.S.A. 85:2444 (1988), or by computerized implementations of these algorithms (for example, GAP, BESTFIT, FASTA, and TFASTA in the Wisconsin Genetics Software Package Release 7.0, Genetics Computer Group, 575 Science Dr., Madison, WI).

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The terms "substantial identity" or "substantial sequence identity" as applied to nucleic acid sequences and as used herein and denote a characteristic of a polynucleotide sequence, wherein the polynucleotide comprises a sequence that has at least 85 percent sequence identity, preferably at least 90 to 95 percent sequence identity, and more preferably at least 99 percent sequence identity as compared to a reference sequence over a comparison window of at least 20 nucleotide positions, frequently over a window of at least 25-50 nucleotides, wherein the percentage of sequence identity is calculated by comparing the reference sequence to the polynucleotide sequence which may include deletions or additions which total 20 percent or less of the reference sequence over the window of comparison. The reference sequence may be a subset of a larger sequence.

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As applied to polypeptides, the terms "substantial identity" or "substantial sequence identity" mean that two peptide sequences, when optimally aligned, such as by the programs GAP or BESTFIT using default gap weights, share at least 80 percent sequence identity, preferably at least 90 percent sequence identity, more preferably at least 95 percent sequence identity or more.

"Percentage amino acid identity" or "percentage amino acid sequence identity" refers to a comparison of the amino acids of two polypeptides which, when optimally aligned, have approximately the designated percentage of the same amino acids. For example, "95% amino acid identity" refers to a comparison of the amino acids of two polypeptides which when optimally aligned have 95% amino acid identity. Preferably, residue positions which are not identical differ by conservative amino acid substitutions. For example, the substitution of amino acids having similar chemical properties such as charge or polarity are not lik by to effect the properties of a protein. Examples include glutamine for asparagine or glutamic acid for aspartic acid.

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The phrase "substantially purified" or "isolated" when referring to a peptide or protein, means a chemical composition which is essentially free of other cellular components. It is preferably in a homogeneous state although it can be in either a dry or aqueous solution. Purity and homogeneity are typically determined using analytical chemistry techniques such as polyacrylamide gel electrophoresis or high performance liquid chromatography. A protein which is the predominant species present in a preparation is substantially purified. Generally, a substantially purified or isolated protein will comprise more than 80% of all macromolecular species present in the preparation. Preferably, the protein is purified to represent greater than 90% of all macromolecular species present. More preferably the protein is purified to greater than 95%, and most preferably the protein is purified to essential homogeneity, wherein other macromolecular species are not detected by conventional techniques.

The phrase "specifically binds to an antibody" or "specifically immunoreactive with", when referring to a protein or peptide, refers to a binding reaction which is determinative of the presence of the protein in the presence of a heterogeneous population of proteins and other biologies. Thus, under designated immunoassay conditions, the specified antibodies bind to a particular protein and do not bind in a significant amount to other proteins present in the sample. Specific binding to an antibody under such conditions may require an antibody that is selected for its specificity for a particular protein. A variety of immunoassay formats may be used to select antibodies specifically immunoreactive with a particular protein. For example, solid-phase ELISA immunoassays are routinely used to select monoclonal antibodies specifically immunoreactive with a protein. See Harlow and Lane (1988) Antibodies, a Laboratory Manual, Cold Spring Harbor Publications, New York, for a description of immunoassay formats and conditions that can be used to determine specific immunoreactivity.

As used herein, "EST" or "Expressed Sequence Tag " refers to a partial DNA or cDNA sequence of about 150 to 500, more preferably about 300, sequential nucleotides of a longer sequence obtained from a genomic or cDNA library prepared from a selected cell, cell type, tissue or tissue type, or organisms which longer sequence corresponds to an mRNA or a gene found in that library. An EST is generally DNA. One or more libraries made from a single tissue type typically provide at least 3000 different (i.e. unique) EST's and potentially the full complement of all possible EST's representing all possible cDNAs, e.g., 50,000 - 100,000 in an animal such as a human. (See, for example, Adams et al. Science 252:1651-1656 (1991)).

"Stringent" as used herein refers to hybridization and wash conditions of 50% formamide at 42°C. Other stringent hybridization conditions may also be selected. Generally, stringent conditions are selected to be about 5° C lower than the thermal melting point (Tm) for the specific sequence at a defined ionic strength and pH. The Tm is the temperature (under defined ionic strength and pH) at which 50% of the target sequence hybridizes to a perfectly matched probe. Typically, stringent conditions will be those in which the salt concentration is at least about 0.02 molar at pH 7 and the temperature is at least about 60°C. As other factors may significantly affect the stringency of hybridization, including, among others, base composition and size of the complementary strands, the presenc of organic solvents and the extent of base mismatching, the combination of parameters is more important than the absolute measure of any one.

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#### B. <u>Transcript Map and New Genes near HH</u>

The instant invention provides a fine structure map of the 1 megabase region surrounding th HFE gene. As part of that map the instant invention provides approximately 250 kb of DNA sequence of which about 235 kb are provided in Figure 8 and eight loci of particular interest corresponding to candidate genes within the 1 megabase region. These loci are useful as genetic and physical markers for further mapping studies. Additionally, the eight cDNA sequences corresponding to those loci are useful, for example, for the isolation of other genes in putative gene families, the identification of homologs from other species, and as probes for diagnostic assays. In particular, isolated nucleic acid sequences of at least 18 nucleotides substantially identical to contiguous nucleotides of a cDNA of the invention are useful as PCR primers. Typically, the PCR primer will be used as part of a pair of primers in a PCR reaction. Isolated nucleic acid sequences preferably comprising about 18-100 nucleotides, more preferably at least 18 nucleotides, substantially identical to contiguous nucleotides in a cDNA of the invention are useful in the design of PCR primers and probes for hybridization assays. Additionally, the proteins encoded by those cDNAs are useful in the generation of antibodies for analysis of gene expression and in diagnostic assays, and in the purification of related proteins.

Thus, in one embodiment of the invention, a 235 kb sequence is provided for the HFE subregion within the 1 megabase region mapped. This sequence can serve as a reference in genetic or physical analysis of deletions, substitutions, and insertions in that region. Additionally, the sequence information provides a resource for the further identification of new genes in that region. Thus, nucleic acid sequences substantially identically to the 235 kb sequence are also included in the scope of this invention.

In a further embodiment of the invention, a family of five genes, BTF1-5, is provided which are related by sequence homology to the milk protein butyrophilin (BT) (Figures 1, 3, and 7). The predicted amino acid sequences of the proteins encoded by these genes are provided in Figure 3. These cDNAs are useful for the identification of further members of the BT family and to study regulation of expression of this family of genes. The proteins encoded by these cDNAs can be useful in the identification and isolation of ligands for the BT protein, and in the generation of agonists or antagonists of BT function. Nucleic acid sequences substantially identically to BTF1-5 and the proteins encoded by them are also included in the scope of this invention, including allelic forms.

In a further embodiment of the invention, a novel gene RoRet is provided, which is related by sequence homology to the 52 kD Ro/SSA Lupus and Sjogren's syndrome autoantigen. This sequence is especially useful in the identification of other genes that may be involved in Lupus or Sjorgen's syndrome. The protein encoded by this cDNA can be useful in the identification and isolation of ligands for the autoantigen, and in the generation of agonists or antagonists of the antigen. Nucleic acid sequences substantially identically to RoRet and the proteins encoded by them are also included in the scope of this invention.

In a further embodiment of the invention, two genes, NPT3 and NPT4, with structural homology to a type 1 sodium transport gene are provided. These cDNAs and the proteins expressed by them ar us full in differentiating the etiology of hypophosphat mia, along with being useful as probes

in the identification and isolation of further members of the gene family. Nucleic acid sequences substantially identically to the NPT1-like sequences and the proteins included by them are also included in the scope of this invention.

## C. Polymorphic Markers

5 The invention provides 397 new polymorphic sites in the region of the HFE gene.

These polymorphisms are listed in Table 1. As described below, these polymorphisms were identified by comparison of the DNA sequence of an affected individual homozygous for the common ancestral HH mutation with that of an unaffected individual disclosed in copending U.S. 08/724,394.

## Table 1. Polymorphic Sites in the HH Region

Base Location	Difference	Base Location	Difference
35-36	AC DEL	19755	G-A
841	T-C ·	19949	C-T
2662-2663	TT DEL	20085	C-T
3767	T-C	20366-20367	AINS
3829	C-G	20463	C-A
4925-4928	TAAA DEL	20841	A-T
5691	C-T	21059	A-T
5839	T-C	21117	A-G
6011	G-A	21837	A-C
6047	C-G	22293	A-C
6231	G-A	22786	C-A
6643	A DEL	23009	G-A
6698	T-C	24143	T-A
7186	T-C	26175	G-C
7273	G-A	26667	C-A
7545-7558	TCACACACCGATTGG DEL	26994	T-C
7672	G DEL	27838	G-T
7933	T-C	27861	T DEL
8746	T-G	28132	G-A
9115	G-A	29100	G-A
9823	G-A	29454-29457	TTTT DEL
10027	G-A	29787	T-G
10214	C-T	29825	A-C
10828	A-G	30009	T-C
10918	C-G	30177	A-G
10955	A-G	30400	A-G
11524	C-A	31059	T-A
11674	A-G	31280	C-T
11955	T-C	31749	C-T
12173-12175	TTT DEL	32040	C-G
13304	G-A	32556-32559	TGTG DEL
13455	G-A	33017	T-G
14416-14417	AINS	33026	T DEL
14998	С-Т	34434	C-T
15564	T-C	35179	A-C
15887	A-G	35695	G-A
15904-15919	CCAAACTGATCTTTGA DEL	35702	G-A
16019	T DEL	35983	A-G
16211	A-T	37411	A-G
17461	A-G	38526	C-T

Base Location	Difference	Base Location	Difference
40431	C-A	72688	C-G
42054-42055	TT DEL	75323-75324	TINS
43783-43784	TTTT INS	75887	G-C
45120	C DEL	77519	T-C
45567	A-C	77749	G-A
46601	A-T	77908	T-C
47255	C-G	78385	C-G
47758	IC-A	78592-78593	AG INS
47994	G-C	80189	T-G
48440	IG-A	80279	T DEL
48650	T-G	80989-80990	AINS
48680	A-G	81193	T-C
50240	C-T	81273	A DEL
50553	G-A	82166	G-A
50586	G-T	83847	T DEL
51322	G-C	84161-84162	CA-GG
	A-G	84533	A-G
51747	C-G	84638	T-G
52474			1
52733	C-A	85526	T-G G-T
52875	G-A	85705	
53631-53637	TTTTTT DEL	86984	T-C
53707	G-A	87655	T-C
54819	A-G	87713 -	A-C
55913	T-C	87892	C-T
56225	A-C	88192	T DEL
56510	T-C_	88528	A-G
56566	G-A	89645	A-T
56618	A-T	89728	A-G
57815	A-G	90088	T-C
58011	T DEL	91193-91194	2209bp INS
58247-58248	TINS	91373	T-C
58926	C-G	91433-91434	AINS
59406	C-G	91747	G-A
59422	G-C	93625	T DEL
60221-60222	AINS	95116-95117	TINS
60656-60657	CA DEL	96315	G-A
61162	IG-A	97981	A-G
61465	G-A	98351	T DEL
61607	A DEL	99249	IC-T
61653	T-C	100094-100095	TINS
	TINS	100647-100648	TTCINS
61794-61795	G-C	100951	IC-T
62061	<del></del>		
62362	T-G	101610	C-G
62732	C-G	102589	IC-T
63364	G-A	103076-103077	TATATATATATA INS
63430-63431	GT INS	103747	T-C
63754	C-T	105638	A-C
63785	A-C	107024	C-T
63870-63871	AINS	107322	C-T
64788	A-G	107858	C-G
64962 °	G-A	109019	A DEL
65891	C-T	109579	T DEL
66675	G-C	110021	C-A
67186-67187	ATT INS	111251	C-A
67746-67747	TTINS	111425	G-A
	T-C	112644	T-A
68259	11.0		
68259 68836			
68259 68836 68976	T-C C-G	113001 113130	G-C C-T

	Base Location	Difference	Base Location	Difference
	114250	A DEL	176222	T-C
	115217	C-G	176524	A-T
	117995	G-A	176684	G-A
	118874	A-G	176815	T-C
5	119470	T-C	177049	T-C
	119646	IG-T	177065	G-T
	120853	C-T	178285	T-C
	121582	G-A	178551-178552	CTTTTTTTTTINS
	123576	A-C	179114-179115	AINS
10	125581	C-T	179260	C-G
	125970	G-T	179281	C-G
	126197	A-G	180023	G-C
	126672	A DEL	180430	T-C
	126672	G-C	180773	T-C
15	128220-128221	AINS	180824	T-C
	132569	C-T	181097	C-T
	133572	A-C	181183	A-T
	134064	T-G	182351	C-T
	136999	G-A	183197	G-A
20	137784	C-T	183623	A-T
	138903	G-A	183653	G-T
	139159-139160	AINS	183657	T-G
	140359	G-A	183795-183796	AINS
	140898	С-Т	184060	G-A
25	141313	CDEL	184993	G-A
	141343	T-C	185918	A-G
	142148	T-C	186036	T-C
	142178	C-A	186506-186507	TAAC INS
30	142433-142434	ATAGA INS	186561-186568	TATTTATT DEL
30	143783 144090	IC-T	186690	G DEL
	144220-144221	A INS	186751 187221	T-A
	144725	IA-C	187260	A-G
	145732-145733	AAAAAAAAAAAAAA INS	187444-187447	A-G CTCT DEL
35	147016-147017	CG DEL	187831-187832	CINS
-	147021	IG-T	188638	G-A
	147536	T-G	188642	C-T
	148936	T-A	189246	T-C
	149061	T-C	190340	A-C
40	154341	A-T	190354	A-G
	154588	G-A	190762	A-G
	155464	G-A	191260	G-T
	158574	C-G	193018-193019	AGAT INS
	160007	IC-T	193147	T-G
45	164348	A-T	193196-193197	CINS
	164499	C-G	193499	C-T
	166677-166678	AAAG INS	193738	C-G
	167389	G-A	193984-193985	ACACACAC INS
50	168506-168507	AGGATGGTCT INS	194064	C-G
50	168515	T-C	194504	A DEL
	169413-169414	AA INS	194734	G-A
•	170300-170301 170491	TTGTTGTTG INS	194890	A-C
	173428	IG-A IT-C	195404	G-A
<b>5</b> 5	173642	IG-A	195693 196205	A-T
-	173948	T-G	190205	G-A C-T
	175330	T-C	197513	C-T
	175836	IT-C	197670	G-A
	176200	G-C	198055	C-A
		_		1.5.1

	Base L cation	Differenc	Bas Location	Difference
	198401	IC-T	215947	C-A
	198692	A-G	216232	A-G
	198780	T DEL	217478	G-A
	199030	T-G	219052	T-C
5	199933	С-Т	219082-219083	ATATATATATATATATATAT INS
	200027	G-A	219314	C-A
	200439	T-A	219327	G-A
	200452	A-G	219560	C-T
	200472-200483	AATAATAATAAT DEL	219660	C-T
10	200559	A-T	219889	G-A
	200745	A-G	220198	G-T
	200919	T-A	220384	G-A
	201816	C-T	220451-220452	CAAAAA INS
	201861-201862	42bp INS	221363	G-A
15	202662	T-C	221645	G-A
	202880	T-C	222119	T-C
	204341	С-Т	222358	A-G
	204768	A-T	222367	A-C
	205284	T-G	222686	A-G
20	207400	C-A	222959	T-C
	208634	T-C	223270-223271	TT DEL
	208718	T DEL	223283	T-C
	208862	A-C	224964	T-C
	209419-209420	TT DEL	225232	A-C
25	209802	G-A	225366-225367	TTTT INS
•	209944	C-G	225416	G-C
	210299	A-G	225486	T-C
	211142	G-A	226088	A-G
	212072	G-A	228421	A-G
30	212146	T-C	230047	G-A
	212379	G-A	230109	G-C
	212637-212639	TCT DEL	230376	C-G
	212696	T-C	230394	A-G
	213042	T-A	231226	A-G
35	214192	A-G	231447	G-A
	214529-214530	TTTTTTTTTINS	231835	A-G
	214549	T-C	232400-232402	AAA DEL
	214795	C-T	232402-232403	G INS
	214908	T-G	232515	T-C
40	214977	A-G	232703	G-T
	215769	C-T	232750	A-G

D6S2238 occurs at base 1. 24d1 occurs at base 41316. D6S2239 occurs at base 84841. D6S2241 occurs at base 235032

## Table 2. Polymorphic Allele Frequencies

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Location	Frequency of ancestral variant in random chromosomes	Frequency of unaffected variant in random chromosomes
232703	53%	47%
231835	53%	47%
230394	85%	15%
230376	25%	75%
230109	53%	47%
225486	45%	55%
225416	75%	25%
220198	43%	57%
219660	58%	42%

	Location	Frequency of ancestral variant in rand m chromosomes	Frequency f unaffected variant in random chromosomes
	219560	53%	47%
	214977	65%	35%
	214908	50%	50%
	214795	24%	76%
5	214549	53%	47%
	214192	65%	35%
	210299	53%	47%
	208862	80%	20%
	208634	48%	52%
10	207400	25%	75%
	205284	50%	50%
	204341	53%	47%
j	202880	58%	42%
	202662	98%	2%
15	200027	25%	75%
	199030	58%	42%
	198692	55%	45%
	198401	55%	45%
	198055	55%	45%
20	195693	60%	40%
	195404	25%	75%
	194890	55%	45%
	175330	53%	47%
	173948	83%	17%
25	173642	55%	45%
	173428	80%	20%
	168515	80%	20%
	160007	18%	82%
į	149061	58%	42%
30	148936	82%	18%
į	147536	100%	0%
	147021	46%	54%
	141343	55%	45%
	140359	55%	45%
35	138903	55%	45%
	132569	81%	19%
ļ	125581	18%	82%
	121582	80%	20%
	120853	18%	82%
40	118874	85%	15%
}	115217	50%	50%
1	113130	40%	60%
1	113001	48%	52%
45	107858	48%	52%
45	103747	50%	50%
	96315	25%	75%
ŀ	91194	80%	20%
}	90088	75%	25%
50	89728 89645	50%	50%
30	88528	50%	50%
-		63%	37%
}	87892 87713	75%	25%
-	87655	60% 50%	40%
55	86984	79%	50%
-	85705	50%	21%
}	85526	50%	50% 50%

	Location	Frequincy of ancestral variant in random chrom somes	Frequency of unaffected variant in random chromos mes
	84638	50%	50%
	84533	50%	50%
	82166	78%	22%
	81193	58%	42%
5	80189	50%	50%
•	78385	80%	20%
	77908	88%	12%
	68976	50%	50%
	68259	51%	49%
10	66675	80%	20%
,,	62732	50%	50%
	62362	40%	60%
	61653	48%	<del></del>
	61465	5%	52% 95%
15	61162	60%	
,5	53707	100%	40%
	52875	50%	0%
	52733	74%	50%
	52474	47%	26%
20	50586	50%	53%
20	50553		50%
	50240	50% 50%	50%
	48680	53%	50%
	48650	63%	47%
25	48440		37%
25		50%	50%
	47255	50%	50%
	46601	53%	47%
	45567	49%	51%
20	41316	5%	95%
30	40431	20%	80%
	38526	23%	77%
	37411	70%	30%
l l	35983	5%	95%

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These polymorphisms provide surrogate markers for use in diagnostic assays to detect the likely presence of the mutations 24d1 and/or 24d2, in preferably 24d1, in homozygotes or heterozygotes. Thus, for example, DNA or RNA from an individual is assessed for the presence or absence of a genotype defined by a polymorphic allele of Table 1, wherein, as a result, the absence of a genotype defined by a polymorphic allele of Table 1 indicates the likely absence of the HFE gene mutation in the genome of the individual and the presence of the genotype indicates the likely presence of the HFE gene mutation in the genome of the individual.

These markers may be used singly, in combination with each other, or with other polymorphic markers (such as those disclosed in co-pending PCT application WO 96/06583) in diagnostic assays for the likely presence of the HFE gene mutation in an individual. For example, any of the markers defined by the polymorphic sites of Table 1 can be used in diagnostic assays in combination with 24d1 or 24d2, or at least one of polymorphisms HHP-1, HHP-19, or HHP-29, or microsatellite repeat alleles 19D9:205; 18B4:235; 1A2:239; 1E4:271; 24E2:245; 2B8:206; 3321-1:98; 4073-1:182; 4440-1:180; 4440-2:139; 731-1:177; 5091-1:148; 3216-1:221; 4072-2:170; 950-1:142;

950-2:164; 950-3:165; 950-4:128; 950-6:151; 950-8:137; 63-1:151; 63-2:113; 63-3:169; 65-1:206; 65-

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2:159; 68-1:167; 241-5:108; 241-29:113; 373-8:151; and 373-29:113, D6S258:199, D6S265:122, D6S105:124; D6S306:238; D6S464:206; and D6S1001:180.

Table 2 lists the frequency of about 100 of the alleles defined by the polymorphic sites of the invention in the general population. As is evident from the Table, certain of these alleles are present rarely in the general population. These polymorphisms are thus preferred as surrogate markers in diagnostic assays for the presence of a mutant HFE allele ("gene mutation") such as 24d1 or 24d2. Preferably, the frequency of the polymorphic allele used in the diagnostic assay in the general population is less than about 50%, more preferably less than about 25%, and most preferably less than about 5%. Thus, of the genotypes defined by the alleles listed in Table 2, polymorphisms occurring at base 35983 and base 61465 of Figure 1 are preferred.

It will be understood by those of skill in the art that because they were identified in an ancestral HH homozygote, the haplotypes defined by the polymorphic sites of Table 1 are predictive of the likely presence of the HFE gene mutation 24d1. Thus, for example, the likelihood of any affected individual having at least two or more of <u>any</u> of the polymorphic alleles defined by Table 1 is greater than that for any unaffected individual. Similarly, the likelihood of any affected individual having at least three or more of <u>any</u> of the polymorphic alleles defined by Table 1 is greater than that for any unaffected individual.

Thus, for example, in a diagnostic assay for the likely presence of the HFE gene mutation in the genome of the individual, DNA or RNA from the individual is assessed for the presence or absence of a haplotype of Table 1, wherein, as a result, the absence of a haplotype of Table 1 indicates the likely absence of the HFE gene mutation in the genome of the individual and the presence of the haplotype indicates the likely presence of the HFE gene mutation in the genome of the individual.

The markers defined by the polymorphic sites of Table 1 are additionally useful as markers for genetic analysis of the inheritance of certain HFE alleles and other genes which occur within the chromosomal region corresponding to the sequence of Figure 9 which include, for example, those disclosed in copending U.S.S.N. 08/724,394.

As the entire nucleotide sequence of the region is provided in Figure 9, it will be evident to those of ordinary skill in the art which sequences to use as primers or probes for detecting each polymorphism of interest. Thus, in some embodiments of the invention, the nucleotide sequences of the invention include at least one oligonucleotide pair selected from the sequence of Figure 9 or its complement for amplification of a polymorphic site of Table 1. Furthermore, in some embodiments of the invention a preferred hybridization probe is an oligonucleotide comprising at least 8 to about 100 consecutive bases from the sequence of Figure 9, or the complement of the sequence, wherein the at least 8 to about 100 consecutive bases includes at least one polymorphic site of Table 1. In some embodiments the polymorphic site is at base 35983 or base 61465.

It will also be appreciated that the nucleic acid sequences of the invention include isolated nucleic acid molecules comprising about 100 consecutive bases to about 235 kb substantially identical to the silvence of Figure 9, wherein the DNA molecule comprises at least one polymorphic

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site of Table 1. Such isolated DNA sequences are useful as primers, probes, or as the component of a kit in diagnostic assays for detecting the likely presence of the HFE gene mutation in an individual.

## D. Nucleic Acid Based Screening

Individuals carrying polymorphic alleles of the invention may be detected at either the DNA, the RNA, or the protein level using a variety of techniques that are well known in the art. The genomic DNA used for the diagnosis may be obtained from body cells, such as those present in peripheral blood, urine, saliva, bucca, surgical specimen, and autopsy specimens. The DNA may be used directly or may be amplified enzymatically *in vitro* through use of PCR (Saiki et al. <u>Science</u> 239:487-491 (1988)) or other *in vitro* amplification methods such as the ligase chain reaction (LCR) (Wu and Wallace <u>Genomics</u> 4:560-569 (1989)), strand displacement amplification (SDA) (Walker et al. <u>Proc. Natl. Acad. Sci. U.S.A.</u> 89:392-396 (1992)), self-sustained sequence replication (3SR) (Fahy et al. <u>PCR Methods Appl.</u> 1:25-33 (1992)), prior to mutation analysis. The methodology for preparing nucleic acids in a form that is suitable for mutation detection is well known in the art.

The detection of polymorphisms in specific DNA sequences, such as in the region of the HFE gene, can be accomplished by a variety of methods including, but not limited to, restrictionfragment-length-polymorphism detection based on allele-specific restriction-endonuclease cleavage (Kan and Dozy Lancet ii:910-912 (1978)), hybridization with allele-specific oligonucleotide probes (Wallace et al. Nucl Acids Res 6:3543-3557 (1978)), including immobilized oligonucleotides (Saiki et al. Proc. Natl. Acad. Sci. U.S.A. 86:6230-6234 (1989)) or oligonucleotide arrays (Maskos and Southern Nucl Acids Res 21:2269-2270 (1993)), allele-specific PCR (Newton et al. Nucl Acids Res 17:2503-2516 (1989)), mismatch-repair detection (MRD) (Faham and Cox Genome Res 5:474-482 (1995)), binding of MutS protein (Wagner et al. Nucl Acids Res 23:3944-3948 (1995), denaturing-gradient gel electrophoresis (DGGE) (Fisher and Lerman et al. Proc. Natl. Acad. Sci. U.S.A. 80:1579-1583 (1983)), single-strand-conformation-polymorphism detection (Orita et al. Genomics 5:874-879 (1983)), RNAase cleavage at mismatched base-pairs (Myers et al. Science 230:1242 (1985)), chemical (Cotton et al. Proc. Natl. Acad. Sci. U.S.A. 85:4397-4401 (1988)) or enzymatic (Youil et al. Proc. Natl. Acad. Sci. U.S.A. 92:87-91 (1995)) cleavage of heteroduplex DNA, methods based on allele specific primer extension (Syvänen et al. <u>Genomics</u> 8:684-692 (1990)), genetic bit analysis (GBA) (Nikiforov et al. <u>Nucl</u> Acids Res 22:4167-4175 (1994)), the oligonucleotide-figation assay (OLA) (Landegren et al. Science 241:1077 (1988)), the allele-specific ligation chain reaction (LCR) (Barrany Proc. Natl. Acad. Sci. U.S.A. 88:189-193 (1991)), gap-LCR (Abravaya et al. Nucl Acids Res 23:675-682 (1995)), radioactive and/or fluorescent DNA sequencing using standard procedures well known in the art, and peptide nucleic acid (PNA) assays (Orum et al., Nucl. Acids Res. 21:5332-5356 (1993); Thiede et al., Nucl. Acids Res. 24:983-984 (1996)).

In addition to the genotypes defined by the polymorphisms of the invention, as described in co-pending PCT application WO 96/35802 published November 14, 1996, genotypes characterized by the presence of the alleles 19D9:205; 18B4:235; 1A2:239; 1E4:271; 24E2:245; 2B8:206; 3321-1:98 (denoted 3321-1:197 therein); 4073-1:182; 4440-1:180; 4440-2:139; 731-1:177; 5091-1:148; 3216-1:221; 4072-2:170 (denoted 4072-2:148 therein); 950-1:142; 950-2:164; 950-3:165; 950-4:128; 950-6:151; 950-8:137; 63-1:151; 63-2:113; 63-3:169; 65-1:206; 65-2:159; 68-1:167; 241-

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5:108; 241-29:113; 373-8:151; and 373-29:113, alleles D6S258:199, D6S265:122, D6S105:124, D6S306:238, D6S464:206; and D6S1001:180, and/or alleles associates with the HHP-1, the HHP-19 or HHP-29 single base-pair polymorphisms can also be used to assist in the identification of an individual whose genome contains 24d1 and/or 24d2. For example, the assessing step can be performed by a process which comprises subjecting the DNA or RNA to amplification using oligonucleotide primers flanking a polymorphism of Table 1, and oligonucleotides flanking 24d1 and/or 24d2, oligonucleotide primers flanking at least one of the base-pair polymorphisms HHP-1, HHP-19, and HHP-29, oligonucleotide primers flanking at least one of the microsatellite repeat alleles, or oligonucleotide primers for any combination of polymorphisms or microsatellite repeat alleles thereof.

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Oligonucleotides useful in diagnostic assays are typically at least 8 consecutive nucleotides in length, and may range upwards of 18 nucleotides in length to greater than 100 or more consecutive nucleotides. Such oligonucleotides can be derived from either the genomic DNA of Figure 8 or 9, or cDNA sequences derived therefrom, or may be synthesized.

Additionally, the proteins encoded by such cDNAs are useful in the generation of antibodies for analysis of gene expression and in diagnostic assays, and in the purification of related proteins.

#### E. **General Methods**

The nucleic acid compositions of this invention, whether RNA, cDNA, genomic DNA, or a hybrid of the various combinations, may be isolated from natural sources, including cloned DNA, or may be synthesized in vitro. The nucleic acids claimed may be present in transformed or transfected whole cells, in a transformed or transfected cell lysate, or in a partially purified or substantially pure form.

Techniques for nucleic acid manipulation of the nucleic acid sequences of the invention such as subcloning nucleic acid sequences encoding polypeptides into expression vectors, labeling probes, DNA hybridization, and the like are described generally in Sambrook et al., Molecular Cloning - a Laboratory Manual (2nd Ed.), Vol. 1-3, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, (1989), which is incorporated herein by reference. This manual is hereinafter referred to as "Sambrook et al."

There are various methods of isolating the nucleic acid sequences of the invention. For example, DNA is isolated from a genomic or cDNA library using labeled oligonucleotide probes having sequences complementary to the sequences disclosed herein. Such probes can be used directly in hybridization assays. Alternatively probes can be designed for use in amplification techniques such as PCR.

To prepare a cDNA library, mRNA is isolated from tissue such as heart or pancreas, preferably a tissue wherein expression of the gene or gene family is likely to occur. cDNA is prepared from the mRNA and ligated into a recombinant vector. The vector is transfected into a recombinant host for propagation, screening and cloning. Methods for making and screening cDNA libraries are well known. See Gubler, U. and Hoffman, B.J. Gene 25:263-269 (1983) and Sambrook et al.

For a genomic library, for example, the DNA is extracted from tissue and either m chanically sh ared or enzymatically digested to yield fragments of about 12-20 kb. The fragments WO 98/14466 PCT/US97/17658

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are then separated by gradient c ntrifugation from und sired sizes and are constructed in bacteriophage lambda vectors. These vectors and phage are packaged *in vitro*, as described in Sambrook, *et al.* Recombinant phage are analyzed by plaque hybridization as described in Benton and Davis, <u>Science</u> 196:180-182 (1977). Colony hybridization is carried out as generally described in M. Grunstein *et al.* <u>Proc. Natl. Acad. Sci. USA.</u> 72:3961-3965 (1975).

DNA of interest is identified in either cDNA or genomic libraries by its ability to hybridize with nucleic acid probes, for example on Southern blots, and these DNA regions are isolated by standard methods familiar to those of skill in the art. See Sambrook, et al.

In PCR techniques, oligonucleotide primers complementary to the two 3' borders of the DNA region to be amplified are synthesized. The polymerase chain reaction is then carried out using the two primers. See <u>PCR Protocols: a Guide to Methods and Applications</u> (Innis, M, Gelfand, D., Sninsky, J. and White, T., eds.), Academic Press, San Diego (1990). Primers can be selected to amplify the entire regions encoding a full-length sequence of interest or to amplify smaller DNA segments as desired.

PCR can be used in a variety of protocols to isolate cDNA's encoding a sequence of interest. In these protocols, appropriate primers and probes for amplifying DNA encoding a sequence of interest are generated from analysis of the DNA sequences listed herein. Once such regions are PCR-amplified, they can be sequenced and oligonucleotide probes can be prepared from sequence obtained.

Oligonucleotides for use as primers or probes are chemically synthesized according to the solid phase phosphoramidite triester method first described by Beaucage, S.L. and Carruthers, M.H., <u>Tetrahedron Lett.</u>, 22(20):1859-1862 (1981) using an automated synthesizer, as described in Needham-VanDevanter, D.R., et al., <u>Nucleic Acids Res.</u>, 12:6159-6168 (1984). Purification of oligonucleotides is by either native acrylamide gel electrophoresis or by anion-exchange HPLC as described in Pearson, J.D. and Regnier, F.E., <u>J. Chrom.</u>, 255:137-149 (1983). The sequence of the synthetic oligonucleotide can be verified using the chemical degradation method of Maxam, A.M. and Gilbert, W., in Grossman, L. and Moldave, D., eds. Academic Press, New York, <u>Methods in Enzymology</u> 65:499-560 (1980).

## 1. Expression

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Once DNA encoding a sequence of interest is isolated and cloned, one can express the encoded proteins in a variety of recombinantly engineered cells. It is expected that those of skill in the art are knowledgeable in the numerous expression systems available for expression of DNA encoding a sequence of interest. No attempt to describe in detail the various methods known for the expression of proteins in prokaryotes or eukaryotes is made here.

In brief summary, the expression of natural or synthetic nucleic acids encoding a sequence of interest will typically be achieved by operably linking the DNA or cDNA to a promoter (which is either constitutive or inducible), followed by incorporation into an expression vector. The vectors can be suitable for replication and integration in either prokaryotes or eukaryotes. Typical expression vectors contain transcription and translation to rminators, initiation sequences, and promotors useful for regulation of the expression of polynucleotide sequence of interest. To obtain

high level xpression of a cloned gene, it is desirable to construct expression plasmids which contain, at the minimum, a strong promoter to direct transcription, a ribosome binding site for translational initiation, and a transcription/translation terminator. The expression vectors may also comprise generic expression cassettes containing at least one independent terminator sequence, sequences permitting replication of the plasmid in both eukaryotes and prokaryotes, i.e., shuttle vectors, and selection markers for both prokaryotic and eukaryotic systems. See Sambrook et al. Examples of expression of ATP-sensitive potassium channel proteins in both prokaryotic and eukaryotic systems are described below.

#### a. Expression in Prokaryotes

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A variety of procaryotic expression systems may be used to express the proteins of the invention. Examples include *E. coli*, *Bacillus*, *Streptomyces*, and the like.

It is preferred to construct expression plasmids which contain, at the minimum, a strong promoter to direct transcription, a ribosome binding site for translational initiation, and a transcription/translation terminator. Examples of regulatory regions suitable for this purpose in *E. coli* are the promoter and operator region of the *E. coli* tryptophan biosynthetic pathway as described by Yanofsky, C., J. Bacteriol. 158:1018-1024 (1984) and the leftward promoter of phage lambda (P\Lambda) as described by Herskowitz, I. and Hagen, D., Ann. Rev. Genet. 14:399-445 (1980). The inclusion of selection markers in DNA vectors transformed in *E. coli* is also useful. Examples of such markers include genes specifying resistance to ampicillin, tetracycline, or chloramphenicol. See Sambrook et al. for details concerning selection markers for use in *E. coli*.

To enhance proper folding of the expressed recombinant protein, during purification from *E. coli*, the expressed protein may first be denatured and then renatured. This can be accomplished by solubilizing the bacterially produced proteins in a chaotropic agent such as guanidine HCl and reducing all the cysteine residues with a reducing agent such as beta-mercaptoethanol. The protein is then renatured, either by slow dialysis or by gel filtration. See U.S. Patent No. 4,511,503.

Detection of the expressed antigen is achieved by methods known in the art as radioimmunoassay, or Western blotting techniques or immunoprecipitation. Purification from *E. coli* can be achieved following procedures such as those described in U.S. Patent No. 4,511,503.

## b. <u>Expression in Eukaryotes</u>

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A variety of eukaryotic expression systems such as yeast, insect cell lines, bird, fish, and mammalian cells, are known to those of skill in the art. As explained briefly below, a sequence of interest may be expressed in these eukaryotic systems.

Synthesis of heterologous proteins in yeast is well known. <u>Methods in Yeast Genetics</u>, Sherman, F., et al., Cold Spring Harbor Laboratory, (1982) is a well recognized work describing the various methods available to produce the protein in yeast.

Suitable vectors usually have expression control sequences, such as promoters, including 3-phosphoglycerate kinase or other glycolytic enzymes, and an origin of replication, termination sequences and the like as desired. For instance, suitable vectors are described in the literature (Botstein, et al., Gene 8:17-24 (1979); Broach, et al., Gene 8:121-133 (1979)).

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Two procedur is are used in transforming yeast cells. In one case, yeast cells are first converted into protoplasts using zymolyase, lyticase or glusulase, followed by addition of DNA and polyethylene glycol (PEG). The PEG-treated protoplasts are then regenerated in a 3% agar medium under selective conditions. Details of this procedure are given in the papers by J.D. Beggs, Nature (London) 275:104-109 (1978); and Hinnen, a., et al., Proc. Natl. Acad. Sci. U.S.A. 75:1929-1933 (1978). The second procedure does not involve removal of the cell wall. Instead the cells are treated with lithium chloride or acetate and PEG and put on selective plates (Ito, H., et al., J. Bact. 153:163-168 (1983)).

The proteins of the invention, once expressed, can be isolated from yeast by lysing the cells and applying standard protein isolation techniques to the lysates. The monitoring of the purification process can be accomplished by using Western blot techniques or radioimmunoassay or other standard immunoassay techniques.

The sequences encoding the proteins of the invention can also be ligated to various expression vectors for use in transforming cell cultures of, for instance, mammalian, insect, bird or fish origin. Illustrative of cell cultures useful for the production of the polypeptides are mammalian cells. Mammalian cell systems often will be in the form of monolayers of cells although mammalian cell suspensions may also be used. A number of suitable host cell lines capable of expressing intact proteins have been developed in the art, and include the HEK293, BHK21, and CHO cell lines, and various human cells such as COS cell lines, HeLa cells, myeloma cell lines, Jurkat cells, etc. Expression vectors for these cells can include expression control sequences, such as an origin of replication, a promoter (e.g., the CMV promoter, a HSV tk promoter or pgk (phosphoglycerate kinase) promoter), an enhancer (Queen et al. Immunol. Rev. 89:49 (1986)), and necessary processing information sites, such as ribosome binding sites, RNA splice sites, polyadenylation sites (e.g., an SV40 large T Ag poly A addition site), and transcriptional terminator sequences. Other animal cells useful for production of ATP-sensitive potassium channel proteins are available, for instance, from the American Type Culture Collection Catalogue of Cell Lines and Hybridomas (7th edition, (1992)).

Appropriate vectors for expressing the proteins of the invention in insect cells are usually derived from the SF9 baculovirus. Suitable insect cell lines include mosquito larvae, silkworm, armyworm, moth and *Drosophila* cell lines such as a Schneider cell line (See Schneider J. Embryol. Exp. Morphol. 27:353-365 (1987).

As indicated above, the vector, e.g., a plasmid, which is used to transform the host cell, preferably contains DNA sequences to initiate transcription and sequences to control the translation of the protein. These sequences are referred to as expression control sequences.

As with yeast, when higher animal host cells are employed, polyadenylation or transcription terminator sequences from known mammalian genes need to be incorporated into the vector. An example of a terminator sequence is the polyadenylation sequence from the bovine growth hormone gene. Sequences for accurate splicing of the transcript may also be included. An example of a splicing sequence is the VP1 intron from SV40 (Sprague, J. et al., J. Virol, 45: 773-781 (1983)).

Additionally, gene sequences to control replication in the host cell may be incorporated into the vector such as those found in bovin papilloma virus type-vectors.

Saveria-Campo, M., 1985, "Bovine Papilloma virus DNA a Eukaryotic Cloning Vector" in <u>DNA Cloning Vol. II a Practical Approach</u> Ed. D.M. Glover, IRL Press, Arlington, Virginia pp. 213-238.

The host cells are competent or rendered competent for transformation by various means. There are several well-known methods of introducing DNA into animal cells. These include: calcium phosphate precipitation, fusion of the recipient cells with bacterial protoplasts containing the DNA, treatment of the recipient cells with liposomes containing the DNA, DEAE dextran, electroporation and micro-injection of the DNA directly into the cells.

The transformed cells are cultured by means well known in the art (Biochemical Methods in Cell Culture and Virology, Kuchler, R.J., Dowden, Hutchinson and Ross, Inc., (1977)). The expressed polypeptides are isolated from cells grown as suspensions or as monolayers. The latter are recovered by well known mechanical, chemical or enzymatic means.

## 2. Purification

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The proteins produced by recombinant DNA technology may be purified by standard techniques well known to those of skill in the art. Recombinantly produced proteins can be directly expressed or expressed as a fusion protein. The protein is then purified by a combination of cell lysis (e.g., sonication) and affinity chromatography. For fusion products, subsequent digestion of the fusion protein with an appropriate proteolytic enzyme releases the desired polypeptide.

The polypeptides of this invention may be purified to substantial purity by standard techniques well known in the art, including selective precipitation with such substances as ammonium sulfate, column chromatography, immunopurification methods, and others. See, for instance, R. Scopes, <u>Protein Purification: Principles and Practice</u>, Springer-Verlag: New York (1982), incorporated herein by reference. For example, in an embodiment, antibodies may be raised to the proteins of the invention as described herein. Cell membranes are isolated from a cell line expressing the recombinant protein, the protein is extracted from the membranes and immunoprecipitated. The proteins may then be further purified by standard protein chemistry techniques as described above.

#### 3. Antibodies

As mentioned above, antibodies can also be used for the screening of polypeptide products encoded by the polymorphic nucleic acids of the invention. In addition, antibodies are useful in a variety of other contexts in accordance with the present invention. Such antibodies can be utilized for the diagnosis of HH and, in certain applications, targeting of affected tissues.

Thus, in accordance with another aspect of the present invention a kit is provided that is suitable for use in screening and assaying for the presence of polypeptide products encoded by the polymorphic nucleic acids of the invention by an immunoassay through use of an antibody which specifically binds to polypeptide products encoded by the polymorphic nucleic acids of the invention in combination with a reagent for detecting the binding of the antibody to the gene product.

Once hybridoma cell lines are prepared, monoclonal antibodies can be made through conventional techniques of priming mice with pristane and interperitoneally injecting such mice with the hybrid cells to enable harvesting of the monoclonal antibodies from ascites fluid.

In connection with synthetic and semi-synthetic antibodies, such terms ar intended to c ver antibody fragments, isotype switched antibodies, humanized antibodies (mouse-human, human-

mouse, and the like), hybrids, antibodies having plural specificities, fully synthetic antibody-like molecules, and the like.

This invention also embraces diagnostic kits for detecting DNA or RNA comprising a polymorphism of Table 1 in tissue or blood samples which comprise nucleic acic, probes as described herein and instructional material. The kit may also contain additional components such as labeled compounds, as described herein, for identification of duplexed nucleic acids.

The following examples are provided to illustrate the invention but not to limit its scope. Other variants of the invention will be readily apparent to one of ordinary skill in the art and are encompassed by the appended claims.

#### 10 F. EXPERIMENTAL EXAMPLES

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#### 1. Megabase transcript map

In these studies direct selection, exon-trapping, and genomic sample sequencing were used to generate a transcript map of a 1 megabase region approximately 8.5 megabases telomeric to HLA-A in the vicinity of HFE. This region 6p21.3 was flanked by the genetic markers D6S2242 and D6S2241. The starting material for these experiments was a 1 megabase YAC labeled y899g1 and a bacterial clone contig of this region (Feder et al. Nature Genetics 13:399-408 (1996)). These techniques and other methods used in the study are outlined below.

#### a. Direct Selection (DS)

Poly A\* RNA from human fetal brain, liver and small intestine (Clontech, Palo Alto, CA) were converted into cDNA using random primers and a Superscript cDNA synthesis kit (Life Technologies, Gaithersburg, MD). The cDNA was digested with Mbo I and ligated to cDNA Mbo I linker-adaptors. Unligated linker-adaptor were removed by passage through cDNA spun columns (Pharmacia, Piscataway, NJ). The 5 ng of each of the ligated cDNAs were amplified using the cDNA Mbo I-S primer (5'-CCTGATGCTCGAGTGAATTC-3'). The amplified products were purified on S-400 spin columns (Pharmacia, Piscataway, NJ), ethanol precipitated and resuspended at 1mg/ml in TE. Gel-purified yac899g1 (Centre d'Etude du Polymorphisme Humain) was processed as described by Morgan et al. (Nucl. Acids Res. 20:5173-5179 (1992)). The cDNAs were mixed in equal molar amounts for a total of 3 mg, and blocked with a mixture of 4 mg Cot-1 DNA (Life Technologies, Gaithersburg, MD), and a cocktail of Sau 3A-digested ribosomal and five different histone DNAs. The blocked cDNAs were hybridized to biotinylated yac899g1 DNA and streptavidin capture was carried out as described by Morgan et al. (ibid). After the second round of selection, the eluted cDNAs were amplified using the cDNA Mbo I-S primer which included a (CUA)4 repeat at the 5' end to facilitate cloning into a version of pSP72 (Promega, Madison, WI) constructed for use with uracil-DNA glycolyase cloning (UDG, Life Technologies, Gaithersburg, MD). Recombinants were transformed in DH5 $\alpha$ , 1000 clones picked into a 96 well format, and clones prepped for DNA sequencing using AGTC boiling 96-well mini-prep system (Advance Genetic Technologies, Gaitherburg, MD).

Four hundred and sixty five clones were sequenced and the resulting data searched by BLAST (Altschul et al. J. Mol. Biol. 215:403-410 (1990)). Those clones representing repetitive, bacterial, yeast, mitochondrial and hist ne sequences were eliminated from futur considerations. The remaining sequences wer then s arched for overlaps and ass mbled into 108 unique DS contigs.

The number of clones per DS contig varied between 1 to 22 with the length of each contig ranging from 250bp to 850 bp. Small sequence-tag-sites PCR assays were developed for each DS contig and two experiments were carried out concomitantly; mapping each DS contig back to the bacterial clone contig of the region and testing for the presence of each DS contig in cDNA libraries. Overall, 86 or 80% of the DS contigs mapped back to the region and were found to be in cDNA libraries. The number of 80% mapping to the region was probably an underestimate of the fidelity of the direct-selection since PCR assays which cross exon-intron boundaries would be expected to fail or give larger size products, thereby being scored negative.

#### b. Exon-Trapping

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CsCl-purified genomic P1 (Genome Systems), BAC (Research Genetics) and PAC (Genome Systems) DNAs were digested with BamHI, Bgl II, Pst I Sac 1 and Xho I and 125 ng of each digest ligated into 500 ng pSPL3 (Church et al. Nature Genetics 6:98-105 (1994)) (Life Technologies, Gaithersburg, MD) digested with the appropriate restriction enzyme and phosphatased with calf intestinal alkaline phosphatase (USB, Cleveland, OH). One tenth of the ligation was used to transform XL1-Blue MRF' cells (Stratagene, La Jolla, CA) by electroporation. Nine tenths of the electroporation was used to inoculate 10 ml of LB + 100μg/ml of carbenicillen and after ovemight growth, DNA was prepared using Qiagen Q-20 tips (Qiagen GmbH, Hilden Germany). The remaining one tenth was plated on LB +100 µg/ml carbenicillen plates to evaluated the efficiency on cloning and to test individual clones for the present of single inserts. COS-7 cells were seed overnight at a density of 1.4 x105/well in 6 well dishes. One µg of DNA was transfected using 6ml of Lipofect-Ace. Cytoplasmic RNA was isolated 48 hr post-transfection. RT-PCR was carried out as described by Church et al. (ibid) using commercially available reagents Life Technologies, Gaithersburg, MD). The resulting CUA-tailed PCR fragments for each restriction digested bacterial clone were pooled and UDG cloned into pSP72-U (a derivative of pSP72). The DNA was transformed in DH5α and the cells plated onto nylon membranes. After overnight growth, duplicates were made and the DNA hybridized to 32P end-labeled oligos designed to detect various background products associated with the pSPL3 vector. One set of filters was hybridized with the following gel-purified oligos in 6X SSC aqueous hybridization solution at 42° C:

vector-vector splicing

5'-CGACCCAGCAACCTGGAGAT-3'

cryptic donor-1021

5'-AGCTCGAGCGGCCGCTGCAG-3'

cryptic donor-1134

5'-AGACCCAACCCACAAGAAG-3'

The filters were washed twice in 6X SSC, 10 mM sodium pyrophosphate (NaPPi) at 60°C, 30 mins.

After overnight autoradiography, non-hybridizing clones were picked and grown in 250 µl of LB + 100µg/ml of carbenicillin in 96 well mini-rack tubes. The samples were analyzed by PCR using the secondary PCR primers supplied in the kit (Life Technologies, Gaithersburg, MD) and those clones with inserts greater than 200 bp were selected for sequencing.

Ninety-six exon traps per bacterial clone were sequenced for a total of 768 reactions and the resulting data analyzed by BLAST. In addition, each potential exon was searched against a database of the 86 DS contigs to eliminat redundant sequences. PCR assays were diveloped for

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each of the potential exons and they were tested for their presence in cDNA libraries. A total of 48 potential exons remained after these screening steps.

#### c. Sample Sequencing

A minimal set of bacterial clones chosen to cover y899g1 were prepped with the Qiagen Maxi-Prep system and purified on CsCl. Ten micrograms of DNA from each bacterial clone was sonicated in a Heat Systems Sonicator XL and end-repaired with Klenow (USB) and T4 polymerase (USB). The sheared fragments were size selected between three to four kilobases on a 0.7% agarose gel and then ligated to BstXl linkers (Invitrogen). The ligations were gel purified on a 0.7% agarose gel and cloned into a pSP72 derivative plasmid vector. The resulting plasmids were transformed into electrocompetent DH5α cells and plated on LB-carbenicillin plates. A sufficient number of colonies was picked to achieve 15-fold clone coverage. The appropriate number of colonies was calculated by the following equation to generate a single-fold sequence coverage: Number of colonies = size of bacterial clone (in kb)/average sequence read length (0.4 kb). These colonies were prepped in the 96-well AGCT system and end-sequenced with oligo MAP1 using standard ABI Dye Terminator protocols. MAP1 was CGTTAGAACGCGGCTACAAT. The MAP1 sequences were screened locally with the BLAST algorithm against all available public databases. All sequence identities were catalogued and cross referenced to the DS and exon-trapped databases.

A total of 3794 end sequence reactions were run to achieve the theoretical 1X coverage. Eighty-five percent of these sequences contained non-bacterial non-vector inserts. An additional 1060 end sequence reactions were run from the opposite end of the cloning vector to augment the sequence coverage and to prepare for contigging across selected regions. BLAST searches to all publicly available databases identified 12 histone genes and 74 unique expressed sequence fragments (ESF). The ESF represent a collection of ESTs and other expressed sequence fragments that were selected due to their sequence identity over a significant portion of genomic DNA. The ESF were cross referenced against the DS and exon-trapped databases to eliminate redundancies. 58 unique ESF remained, representing 39 distinct clones. Included in these ESF are 5 sequences homologous to histone genes.

Table 3. EST's found by Sample Sequencing Large Insert Bacterial Clones

30	Clone name	Bacterial clone	Homology 5' blastx	Homology 3' blastx	Poly A+ signal	Genomic poly (A) <sub>as</sub>	cDNA Homology
	EST03556	pc157c3	na²	none <sup>3</sup>	+	-	cDNA 28
	ym33f11	pc157c3	ZNF	na	па	na	
	EST04698	pc157c3	na	NSH <sup>4</sup>	+	•	
	EST04812	pc157c3	na	NSH	-	•	
35	уь89ь08	pc157c3	NSH	na	na	na	
	yd88g11	pc157c3	na	nsh	+	•	
	yj49b01	pc157c3	NSH	na	na	na	
	yv81d05	pc157c3	HG17 Human	NSH	+	-	cDNA 30
	yg57h09	p196e20	BUTYBOVIN	NSH	+	-	cDNA 21
40	yq23d08	p196e20	BUTYBOVIN	NSH	+	•	cDNA 21

30	Clone name	Bacterial clone	Homology 5' blastx	Homology 3' blastx	Poly A+ signal¹	Genomic poly (A) <sub>os</sub>	cDNA Homology
	yo65f06	p196e20	NSH	na	na	па	cDNA 29
	yv88c09	p196e20	BUTYBOVIN	na	na	na	cDNA 29
	yd17d06	p196e20	NSH	na	na	na	cDNA 23
•	ye25g03	p196e20	BUTYBOVIN	NSH	na	na	cDNA 44
5	ys04h08	pc45p21	NSH	NSH	+	-	cDNA 44
	yn01c05	p196e20	BUTYBOVIN	па	па	na	cDNA 32
	YG78F10	PC45P21	NSH	NSH	na	na	
	yh54f11	p196e20	none	NSH	-	-	
	ys05b08	pc157c3	NSH	Alu		+	
10	yb12h11	b132a12	NSH	Histone H3.1	-	•	
	HSC2EE082	b132a12	na	NSH	+	•	
	HUM160h11b	b132a12	none	na .	na	na	
	yg04f09	b132b12	Line element	Alu	-	+	
	yd37d11	b132a12	NSH	Alu	-	+	
15	ym29g03	b132a12	Histone H2A	NSH	+	•	cDNA 37
	yi77b02	b132a12	NSH	NSH	-	-	cDNA 37
	yh76b05	b132a12	NSH	Alu	-	•	
	yu98e02	b132a12	NSH	Alue	•	+	
	yd72h12	b132a12	Alu	NSH	+	+	
20	yd 19d03	pc222k22	Histone H2B.1	NSH	+	•	
	ye98g01	b132a12	NSH	NSH	+	-	cDNA
	yi61f07	b132a12	NSH	NSH		+	
	ESTO5340	b3e17	na	Alu	•	+	
	yd35d05	pc222k22	NSH	NSH	-	+	
25	yc52a05	pc75L14	NSH	na	па	na	
	yd84a05	pc75L14	none	none	-	?5	-
	уг42а05	pc75L14	NaPi transport	none	+		cDNA 22B
	yd83h08	b20h20	NSH	none	+	•	
	ye38c09	b20h20	NSH	Aiu	•	+	
30	yp74c05	b20h20	NaPi transport	Alu	?6	na	
	Bracketed area is	the critical regi	on				
	1 Signal of	ATAAA or AT	ГАА		4	No Significant I	Homologies
	2 Not avail	able			5	3' splice that is	-
35	3 "NONE"		6	Poor EST seque	<u> </u>		

## d. cDNA library screening

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Superscript plasmid cDNA libraries, brain, liver and testis, were purchased from Life Technologies, Gaithersburg, MD. Colonies were plated on Hybond N filters (Am rsham) using

standard techniques. Insert probes from DS, exons and EST (I.M.A.G.E. clones; Genome Systems) were all isolated by PCR followed by purification in low-melting point agarose gels (Seak m). Th DNAs were labeled in gel using the Prime-it II kit (Stratagene, La Jolla, CA). Small exon probes were labeled using their respective STS PCR primers instead of random primers. Up to 5 different probes were pooled in a hybridization. Filters were hybridized in duplicate using standard techniques. Putative positives were screened by PCR using the probe's STSs to identify clones. Inserts from positive clones were subcloned in pSP72 and sequenced.

#### e. Northern blots and RT-PCR analysis

Multiple tissue northern blots were purchased from Clontech and hybridized according the manufacturer's instructions. RT-PCR was carried out on random primed first strand cDNA made from poly A+ RNA (Clontech) using AmpliTaq Gold (Perkin-Elmer). Control reactions were performed on RNA samples processed in the absence of reverse transcriptase to control for genomic DNA contamination.

#### f. Genomic Sequencing

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The MAP1 sequences from the bacterial clones b132a2, 222K22, and 75L14 were assembled into contigs with the Staden package (available from Roger Staden, MRC). A minimal set of 3 kb clones was selected for sequencing with oligo labeled MAP2 that sits on the opposite end of the plasmid vector. The sequence of MAP2 was GCCGATTCATTAATGCAGGT. The MAP2 sequences were entered into the Staden database in conjunction with the MAP1 sequences to generate a tiling path of 3 kb clones across the region. These sequences were also screened with the BLAST algorithm and all novel sequence identities were noted. The plasmid 3 kb libraries were concurrently transformed in 96 well format into pox38UR (available from C. Martin, Lawrence Berkeley Laboratories). The transformants were subsequently mated with JGM (Strathman et al. P.N.A.S. 88:1247-1250 (1991) in 96 well format. All matings of the 3 kb clones within the tiling path were streaked on LB-carbenicillin-kanamycin plates and a random selection of 12 colonies per 3 kb clone was prepped in the AGCT system. The oligos -21: CTGTAAAACGACGGCCAGTC, and REV: GCAGGAAACAGCTATGACC were used to sequence off both ends of the transposon. Each 3 kb clone was assembled in conjunction with the end sequence information from all bacterial clones to generate complete sequence across the region. The genomic sequence was analyzed with the BLAST nucleotide and protein homology algorithms and the GRAIL 1.2 software to identify novel open reading frames (ORF) for gene finding.

## g. Discussion

A compilation of 174 ESF led to the construction of an expressed sequence map of the region that served as the framework for the isolation of full-length cDNAs (Figure 1). (The map shows the subset of ESF that were actually mapped). Probes were developed for 82 best ESFs which appeared to be derived from the coding portions of cDNAs and the appropriate cDNA libraries were screened. This led to the isolation of 19 cDNAs, 17 of which represented novel sequences. 70 of the 174 ESF were included in the cDNAs isolated (40%). 36 probes failed to produce any clon's even after repeated screening of several libraries. 51 ESF which were not accounted for in the cDNAs

cloned were not used in any screen. Therefore, it is possible that some additional genes within this 1 megabase region may have escaped detection.

A list of these cDNAs cloned and a comparison of the methods used to find them is presented in Table 4. Direct selection found 14 out of the 18 cDNAs contained within the boundaries of the YAC used in the experiment. Exon trapping found 15 out of the 19 cDNAs contained within the boundaries of the large insert bacterial clone contig. Sample sequencing identified 11 genes that had corresponding ESTs in the public database.

Table 4. Comparison of gene finding methods

10		mon or Belie Inio	mg memous			
	Bacterial Clone	CDNA#	Homology	EST	DS	Exon Trap
	157c	28	zinc finger	EST03556	2	1
	157c3	30	nonhistone	yv81d05	1	none
				yvh07a10	•	
	157c3	46	ORF	yd88g11	1	
15	157c3	20	ВТ	none	none	3
	p18696	21	BTF1	yn01G5	4	5
				yg23d08		-
				yg57h09		
				yu15h03		
	45p21	32	BTF2	yg78f10	7	3
			_	yn01c05		•
	45p21	29	BTF3	ye25g03	2	9
				yo65f06		
	45p21	23	BTF4	yd17d06	4	6
20	45p21	44	BTF5	. ys04h08	2	4
	3e17	41	genomic?	none	none	1
	132a2	43	genomic?	none	none	3
	132a2	36	genomic?	none	1	none
	132a2	37	histone 2A	ym29g03	3	none
				yh87a03		
25	75114	24	MHC class 1	ye98g01	1	2
	132a2	39	genomic?	none	none	4
	132a2	27	Ro/SSA	none	3	4
	132a2	22B	NPT1-like	ут42а05	1	7
				yf09g06		
	20h20	22E	NPT1-like	none	2	5
30	20h20	NPT1	NPT1	yp74c05	N/A	3

As a final approach, a tiling path with verlapping end sequences from the sample sequence database was generated. Each 3 kb clone within the path was shotgun-sequenced using transposable elements as platforms for dual end sequencing. Thes individual clones were assembled in conjunction with the end sequences from all bacterial clones in the region. The resulting sequence (Figure 2) was analyzed systematically with BLAST homology searches and the Grail 1.2 program to identify novel open reading frames (ORF) and other gene-like structures. The BLAST homology searches did not produce any probes that had not already been identified by sample sequencing. Grail predicted exons for all the genes in the region, but was only able assemble the histones into any representative form. A detailed analysis of BLAST homology searches to protein databases identified an enticing homology to a zinc alpha 2 glycoprotein approximately 25 kb upstream of HFE, but the lack of a substantial ORF and the presence of a stop codon suggested that it was a pseudogene. Figure 2 shows the positions, the exon and intron structures, and the relative orientation of transcription of novel genes within this region. Also shown are the positions and transcriptional orientations of the histone genes. A total of 12 histone genes were identified in this study.

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In an effort to account for the ESTs that did not associate with the characterized genes in the 250 kb region, the genomic sequence around the putative 3' ends were examined for polyadenylation signals to determine whether certain EST sequences may have originated from genomic DNA contamination in the normalized cDNA libraries used in EST generation. The positions of the 14 ESTs found in this region are indicated in Figure 2 to show those associated with the cDNAs cloned and those which did not associate with genomic DNA of obvious coding potential. Four ESTs corresponded to 3 of the 4 cDNAs cloned from the region (Table 2). One EST encoded a histone H2B.1 gene and another was a repetitive element. Of the remaining 8, 6 EST clones were used as probes of cDNA libraries with negative results. Those sequences representing putative 3' ends of cDNA were searched for the presence of poly (A)+ addition signals. Five of the 13 ESTs which had 3' end sequence, had the sequence ATAAA or ATTAA. Five of the remaining 8 ESTs that did not have a poly (A)+ addition signal had genomic encoded stretches of poly (A) near the end of EST sequence and, therefore, may have been created by oligo d(T) priming of contaminating genomic DNA. This analysis was expanded to include all ESTs in the large-insert bacterial contigs with definitive 3' ends. Of the remaining 26, 15 had 3' end sequence and, of these, 8 had poly (A)+ addition signals. Five of these 8 ESTs were associated with the cloned cDNAs. Of the remaining 7 which did not have poly (A)+ addition signals, 4 had genomic encoded stretches of poly (A).

#### i. Butyrophilin gene family

The human homolog of the bovine butyrophilin gene (BT) was cloned and mapped to approximately 480 kb centromeric to HFE (Figure 1). BT is a transmembrane protein of unknown function which constitutes 40% of the total protein associated with the fat globule of bovine milk (Jack et al. J. Biol. Chem. 265:14481-14486 (1990)). A human homolog of BT has recently been cloned by Tayloer et al. (Biochem Biophys Acta 1306:1-4 (1996)). The results in this study indicated that BT is a member of a gene family with at least five other members of the family residing in this region (Figure 1). A comparison of the se proteins is shown in Figure 3. The proteins were aligned based on their discending order of relatedness and to minimized gaps in the sequence. Each of the five proteins

display varying degrees of homology to BT. BTF1 (cDNA 21), BTF2 (cDNA 32), BTF5 (cDNA 44), and BTF3 (cDNA 29) are 45%, 48%, 46%, and 49%, identical to BT, whereas BTF4 (cDNA 23), which is more similar to BTF3 (cDNA 29), is only 26% identical. This low degree of identity to BT is largely due to a truncation at the carboxyl terminus of the protein. The BTF family falls into two groups: BTF1 and 2 which are more related to each other than to BT or the other BTF members, and BTF5, 3 and 4, which appear to have a common evolutionary origin. The order of these genes on the chromosome suggests that the BT gene has duplicated two times, giving rise to BTF1 and BTF5. Subsequently, it appears likely these two genes experienced further duplication events to give rise to the other members in their groups.

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The three major components of BT, the B-G immunoglobulin superfamily domain (containing the V consensus sequence) (Miller et al. Proc. Natt. Acad. Sci. U.S.A. 88:4377-4381 (1991)), the transmembrane region, and the B30-2 exon are found in all of these proteins (with the exception of BTF4 (cDNA 23) which lacks the B30-2 exon by virtue of the carboxyl terminal truncation). The exon B30-2 is a previously noted feature of the MHC class 1 region found approximately 200 kb centromeric to the HLA-A gene (Vernet et al., J. Mol. Evol. 37:600-612 (1993)). In addition this exon is found in several genes of diverse function telomeric to HLA-A namely MOG (approximately 200 kb) and RFP (approximately 1 megabase) (Amadou et al. Genomics 26:9-20 (1995)).

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The levels of the BTF mRNA were analyzed by northern blot analysis (Figure 4A). The expression of the BTF genes fell into two patterns. BTF1 and BTF2 were expressed as a single major transcript of 2.9 kb and one minor transcript of 5.0 kb. These genes were expressed at high levels in all the tissues tested with the exception of the kidney where the expression level was less. The two genes are 90% identical at the DNA sequence level, therefore, it is possible that the signal observed on the northerns was the result of cross-hybridization and only one of the two genes was actually expressed. To address this possibility RT-PCR experiments were carried out on a panel of different tissues in order to detect possible tissue dependent expression that would suggest that both genes are expressed. Identical, and thus equivocal, results were obtained with both BTF1 and BTF2 amplification (Figure 4B).

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The second group of genes, BTF3-5, are expressed as three (BTF5) (Figure 4A) and two (BTF3 and 4) transcripts ranging from 4.0 to 3.3 kb. BTF5 is expressed at moderate levels in all tissues tested with the exception of the kidney where the expression level is less. RT-PCR experiments showed that mRNA from the BTF5 gene can be found in all tissues tested, including the kidney (Figure 4B). Identical results were obtained with primers from the other genes of this group (data not shown). These genes are also 90% identical to each other at the DNA sequence level (but only 58% identical to BTF1 and 2), hence like BTF1 and BTF2, cross-hybridization could account for the similarity in size and patterns on the northern blots and RT-PCR. This might be particularly true for BTF4 which lacks the B30-2 exon but still hybridizes to larger size transcripts like BTF5 and BTF3.

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## ii. A gene with similarity to 52 kD Ro/SSA auto-antigen

Located approximately 120 kb telement to the HFE gen is a gene, RoRet, that has 58% amino acid similarity to the 52 kD Ro/SSA protein, an auto-antigen of unknown function that is frequently recognized by antibodies in patients with systemic lupus and Sjogren's syndrome (Anderson

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et al. Lancet 2:456-560 (1961); Clark et al. J. Immunol. 102:117-122 (1969)) (Figures 1 and 2). Alignment of the predicted amin—acid sequence of this cDNA with that of 52 kD Ro/SSA indicated two features associated with the 52 kD Ro/SSA protein: a putative DNA binding cysteine rich motif (C-X-(I,V)-C-X(11-30)-C-X-H-X-(F,I,L)-C-X(2)-C-(I,L,M)-X(10-18)-C-P-X-C) found at the N terminus (Freemont et al. Cell 64: 483-484 (1991)) and the B30-2 exon found near the carboxyl terminus, are both conserved in RoRet (Figure 5). Northern blot analysis indicated the RoRet gene was expressed as two major transcripts of 2.8 and 2.2 kb and two minor transcripts of 7.1 and 4.4 kb in all of the tissues on the blot at levels reflective of the RNA amounts as determined by β-a ±tin probing (Figure 6A). Using RT-PCR, expression can also be detected in small intestine, kidney liver, and spleen (Figure 6B).

# iii. Two genes with homology to a sodium phosphate transporter

A cDNA for a sodium phosphate transport protein (NPT1) was previously cloned and mapped to 6p21.3 using a somatic cell hybrid panel (Chong et al. Genomics 18:355-359 (1993)). NPT1 maps 320 kb telomeric to the HFE gene (Figures 1 and 2). Two additional cDNAs were cloned which show appreciable homology to NPT1 (Figure 5). These genes, NPT3 and NPT4, mapped 1.5 megabases and 1.3 megabases centromeric to the NPT1 gene (Figure 1). Like NPT1, the gene products of NPT3 and NPT4 were extremely hydrophobic, which may reflect a membrane location. Both proteins gave hydrophilicity profiles which were indistinguishable from NPT1 in this study (data not shown). Northern blot analysis indicated that the two genes have different patterns of expression (Figure 6C). NPT3 was expressed at high levels as a 7.2 kb transcript predominately in muscle and heart. Lesser amount of the mRNA were also found in brain, placenta, lung, liver and pancreas. RT-PCR analysis indicated that expression of the proper size PCR fragment for NPT3 was clearly absent in fetal brain, bone marrow and small intestine (Figure 6D). A smaller size fragment was detectable in all tissues with the exception of the liver, which may represent evidence for alternative splicing. Although expression was apparently absent from the kidney by northern blot analysis, it was detectable by RT-PCR. Expression was also noted in the mammary gland, spleen and testis. NPT4, on the other hand, was expressed only in the liver and the kidney as a smear of transcripts approximately 2.6 - 1.7 kb (Figure 6C). RT-PCR confirmed these results, although a small amount of the proper size PCR fragment was also found in the small intestine and testis (Figure 6D). Other tissues showed amplification, but the fragments were of larger and smaller size than that produced by the cDNA 22E positive control. Hence, these two genes which apparently have the structural characteristics of a sodium phosphate transporter, appeared to be under the control of different regulatory mechanism that lead to differential patterns of expression.

## 2. Sequencing of 235 kb from a Homozygous Ancestral (Affected) Individual

In these studies the entire genomic sequence was determined from an HH affected individual for a region corresponding to a 235,033 bp region surrounding the HFE gene between the flanking markers D6S2238 and D6S2241. The sequence was derived from a human lymphoblastoid cell line, HC14, that is homozygous for the ancestral HH mutation and region. The sequence from the ancestral chromosome (Figure 9) was compared to the sequence of the region in an unaffected individual (Figure 8) disclosed in copending U.S.S.N. 08/724,394 to identify polymorphic sites. A

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subset of the polymorphic alleles so defined were further studied to determine their frequency in a collection of random individuals.

The cell line HC14 was deposited with the ATCC om June 25, 1997, and is designated ATCC CRL-12371.

## a. Cosmid Library Screening

The strategy and methodology for sequencing the genomic DNA for the affected individual was essentially as described in copending U.S.S.N. 08/724,394, hereby incorporated by reference in its entirety. Basically, a cosmid library was constructed using high molecular weight DNA from HC14 cells. The library was constructed in the supercos vector (Stratagene, La Jolla, CA). Colonies were replicated onto Biotrans nylon filters (ICN) using standard techniques. Probes from genomic subclones used in the generation of the sequence of the unaffected sequence disclosed in 08/724,394 were isolated by gel electrophoresis and electroporation. Subclones were chosen at a spacing of approximately 20 kb throughout the 235 kb region. The DNA was labeled by incorporation of 32P dCTP by the random primer labeling approach. Positively hybridizing clones were isolated to purity by a secondary screening step. Cosmid insert ends were sequenced to determine whether full coverage had been obtained, and which clones formed a minimal path of cosmids through the 235 kb region.

#### b. Sample Sequencing

A minimal set of cosmid clones chosen to cover the 235 kb region were prepped with the Qiagen Maxi-Prep system. Ten micrograms of DNA from each cosmid preparation were sonicated in a Heat Systems Sonicator XL and end-repaired with Klenow (USB) and T4 DNA polymerase (USB). The sheared fragments were size selected between three to four kilobases on a 0.7% agarose gel and then ligated to BstXl linkers (Invitrogen). The ligations were gel purified on a 0.7% agarose gel and cloned into a pSP72 derivative plasmid vector. The resulting plasmids were transformed into electrocompetent DH5α cells and plated on LB-carbenicillin plates. A sufficient number of colonies was picked to achieve 15-fold clone coverage. The appropriate number of colonies was calculated by the following equation to generate a single-fold sequence coverage: Number of colonies = size of bacterial clone (in kb)/average sequence read length (0.4 kb). These colonies were prepped in the 96-well Qiagen REAL, and the 5' to 3' DNA Prep Kit, and AGCT end-sequenced with oligo MAP1 using standard ABI Dye Terminator protocols. MAP1 was CGTTAGAACGCGGCTACAAT.

## c. Genomic Sequencing

The MAP1 sequences from the cosmid clones HC182, HC187, HC189, HC195, HC199, HC200, HC201, HC206, HC207, and HC212 were assembled into contigs with the Staden package (available from Roger Staden, MRC). A minimal set of 3 kb clones was selected for sequencing with oligo labeled MAP2 that sits on the opposite end of the plasmid vector. The sequence of MAP2 was GCCGATTCATTAATGCAGGT. The MAP2 sequences were entered into the Staden database in conjunction with the MAP1 sequences to generate a tiling path of 3 kb clones across the region. The plasmid 3 kb libraries w re concurrently transf rmed in 98 well format into pox38UR (available from C. Martin, Lawrence Berkeley Laboratories). The transformants w re subsequently mated with JGM (Strathman et al. P.N.A.S. 88:1247-1250 (1991) in 96 well format. All matings of the

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3 kb clones within the tiling path were streak d on LB-carbenicillin-kanamycin plates and a random selection of 12 colonies per 3 kb clone was prepped in the AGCT system. The oligos -21: CTGTAAAACGACGCCAGTC, and REV: GCAGGAAACAGCTATGACC were used to sequence off both ends of the transposon. Each 3 kb clone was assembled in conjunction with the end sequence information from all cosmid clones in the region.

In some regions, the coverage of the genomic sequence by cosmids was incomplete. Any gaps in the sequence were filled by using standard PCR techniques to amplify genomic DNA in those regions and standard ABI dye terminator chemistry to sequence the amplification products.

#### d. Identification of Polymorphic Sites

The assembled sequence of the cosmid clones in connection with the PCR amplified genomic DNA was compared to the genomic sequence of the unaffected individual using the FASTA algorithm. Numeric values were assigned to the sequenced regions of 1 to 235,303, wherein base 1 refers to the first C in the CA repeat of D6S2238 and base 235,303 is the last T in the GT repeat of D6S2241 of the <u>unaffected</u> sequence (Figure 8). Table 1 lists the differences between the two compared sequences. Note that previously disclosed (Feder et al., <u>Nature Genetics</u> 13:399-408 (1996)) polymorphic sites D6S2238 (base 1), D6S2241 (base 235,032), 24d1 (base 41316), and D6S2239 (base 84841) are not included in the list of new polymorphisms, although they are provided for reference in a footnote to the Table and were observed in the ancestral sequence. In the Table, a single base change such as C-T refers to a C in the unaffected sequence at the indicated base position that occurred as a T in the corresponding position in the affected sequence. Similarly, an insertion of one or more bases, such as TTT in the affected sequence, is represented as "TTT INS" between the

e. Characterization of Rare Polymorphisms

In this study about 100 of the polymorphisms of Table 1 were arbitrarily chosen for further characterization. Allele frequencies in the general population were estimated by OLA analysis using a population of random DNAs (the "CEPH" collection, J. Dausset et al., <u>Genomics</u> 6(3):575-577 (1990)). These results are provided in Table 2.

indicated bases of the unaffected sequence. A deletion of one or more bases occurring in the affected sequence, such as AAA DEL, is represented as the deletion of the indicated bases in the unaffected

One single base pair difference, occurring at base 35983 and designated C182.1G7T/C (an A to G change on the opposite strand) was present in the ancestral chromosome and rare in the random DNAs. This change occurred in a noncoding region of the hemochromatosis gene near exon 7 approximately 5.3 kb from the 24d1 (Cys282Tyr) mutation. OLA was used to genotype 90 hemochromatosis patients for the C182.1G7T/C base pair change. The frequency for C occurring at this position in the patients was 79.4% as compared to 5% in the random DNAs. Eighty-five of the 90 patients assayed contained identical 24d1 and C182.1G7T/C genotypes. Four of the remaining 5 patients were homozygous at 24d1 and heterozygous at C182.1G7T/C; one was heterozygous at 24d1 and homozygous at C182.1G7T/C. The primers used for this analysis were as follows.

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sequence.

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PCR primers for detection:

182.1G7.F 5'-GCATCAGCGATTAACTTCTAC -3'

182.1G7.R 5'-TTGCATTGTGGTGAAATCAGGG -3'

For the detection assay, the biotinylated primers used were as follows.

182.1G7.C 5' (b)CTGAGTAATTGTTTAAGGTGC -3'

182.1G7.T 5' (b)CTGAGTAATTGTTTAAGGTGT -3'

The phosphorylated digoxigenin-labeled primer used was:

182.1G7.D 5' (p)AGAAGAGATAGATATGGTGG -3'

A further rare single base pair change was detected at 61,465bp. The inheritance pattern of this polymorphism, C195.1H5C/T (a G to A change on the opposite strand), is identical to that of 24d1. The frequency of T occurring at that position (C195.1H5T) observed in a set of 76 patients was 78.5% as compared to 5% in random individuals.

15 PCR primers for detection:

1951H5.3F 5'-GAATGTGACCGTCCCATGAG-3'

1951H5.3R 5'-CAACTGAATATGCAGAAAAAAGTACACC-3'

For the detection assay, the biotinylated primers used were:

1951H5.3.4 5' (b)AGTAGCTGGGACTCACGGTGT-3'

1957H5.3.5 5' (b)AGTAGCTGGGACTCACGGTGC-3'

The phosphorylated digoxigenin-labeled primer used was:

1951H5.3.6 5' (p)GCGCCACCACTCCCAGCTCAT-3'

These rare alleles are thus preferred surrogate markers for 24d1 and are especially useful in screening assays for the likely presence of 24d1 and/or 24d2.

All publications, patents, and patent applications cited herein are hereby incorporated by reference in their entirety.

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#### WHAT IS CLAIMED IS:

1	1.	An oligonucleotide comprising at least 8 to about 100 consecutive bases from the
2	sequence of F	igure 9, or the complement of the sequence, wherein the at least 8 to about 100
3	•	ases includes at least one polymorphic site of Table 1.
1	2.	The oligonucleotide of claim 1, wherein the polymorphic site is selected from the
2	group consistir	ng of base 35983 or base 61465.
1	3.	An oligonucleotide pair selected from the sequence of Figure 9 or its complement for
2	amplification of	f a polymorphic site of Table 1.
1	4.	An isolated nucleic acid molecule comprising about 100 consecutive bases to about
2	235 kb substa	ntially identical to the sequence of Figure 9, wherein the DNA molecule comprises at
3	least one poly	morphic site of Table 1.
1	5.	The isolated nucleic acid molecule of claim 4, wherein the polymorphic site is selected
2	from the grou	p consisting of base 35983 or base 61465.
1	6.	The isolated nucleic acid molecule of claim 4, wherein the nucleic acid is selected
2	from the grou	p consisting of cDNA, RNA, or genomic DNA.
1	7.	A polypeptide encoded by the nucleic acid molecule of claim 4.
1	<b>.</b> 8.	An antibody which specifically recognizes the polypeptide of claim 7.
1	9.	A method to determine the presence or absence of the common hereditary
2	hemochroma	tosis (HFE) gene mutation in an individual comprising:
3		providing DNA or RNA from the individual; and
4		assessing the DNA or RNA for the presence or absence of a haplotype of Table 1,
5	wher	ein, as a result, the absence of a haplotype of Table 1 indicates the likely absence of the
6	HFE gene m	utation in the genome of the individual and the presence of the haplotype indicates the
7	likely present	ce of the HFE gene mutation in the genome of the individual.
1	10.	The method of claim 9, wherein the method further comprises assessing the RNA or
2		presence of at least one of the polymorphisms 24d1, 24d2, HHP-1, HHP-19, or HHP-29;
3		lite repeat alleles 19D9:205, 18B4:235, 1A2:239, 1E4:271, 24E2:245, 2B8:206, 3321-
4	1:98, 4073-1	:182, 4440-1:180, 4440-2:139, 731-1:177, 5091-1:148, 3216-1:221, 4072-2:170, 950-

1:142, 950-2:164, 950-3:165, 950-4:128, 950-6:151, 950-8:137, 63-1:151, 63-2:113, 63-3:169, 65-

6	1:206, 65-2:159	9, 68-1:167, 241-5:108, 241-29:113, 373-8:151, 373-29:113, D6S258:199, D6S255.122,
7	D6S105:124, D	6S306:238, D6S464:206, or D6S1001:180.
1	11.	The method of claim 9, wherein the haplotype comprises at least two polymorphic
2	sites of Table 1	
	42	The method of claim 11, wherein one of the at least two polymorphic sites of Table 1
1 2	12. is at base 3598	
		The method of claim 11, wherein the haplotype comprises at least three polymorphic
1 2	13. sites of Table 1	
		A method to determine the presence or absence of the common hereditary
1	14.	
2	hemochromate	osis (HFE) gene mutation in an individual comprising:
3		providing DNA or RNA from the individual; and assessing the DNA or RNA for the presence or absence of a genotype defined by a
4.		
5		llele of Table 1, in, as a result, the absence of a genotype defined by a polymorphic allele of Table 1
6		kely absence of the HFE gene mutation in the genome of the individual and the
7	indicates the II	e genotype indicates the likely presence of the HFE gene mutation in the genome of the
8 9	presence of the individual.	e genotype indicates the likely presence of the Fin E gene material and general
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1	15.	The method of claim 15, wherein the polymorphic allele occurs in less than about 50%
2	of a random p	opulation of individuals.
1	16.	The method of claim 15, wherein the polymorphic allele occurs in less than about 25%
2	of a random p	opulation of individuals.
1	17.	The method of claim 15, wherein the polymorphic allele occurs in less than about 5%
2		opulation of individuals.
1	18.	The method of claim 15, wherein the genotype is C182.1G7C or C195.1H5T.
1	19.	A kit comprising one or more oligonucleotides of claim 1.
1	20.	A kit comprising at least one oligonucleotide pair of claim 3.
1	21.	A cultur of lymphoblastoid cells having the designation ATCC CRL-12371.

1 2	BTF1.	22.	An isolated nucleic acid sequence comprising a sequence substantially identical to
1		23.	The isolated nucleic acid sequence of claim 23, wherein the nucleic acid is cDNA.
1		24.	The polypeptide encoded by the isolated nucleic acid sequence of claim 23.
1		25.	A vector comprising the nucleic acid sequence of claim 23.
1		26.	A host cell stably transfected with the nucleic acid sequence of claim 23.
1		27.	An antibody that is specifically immunoreactive with the polypeptide of claim 24.
1 2	BTF2.	28.	An isolated nucleic acid sequence comprising a sequence substantially identical to
1		29.	The isolated nucleic acid sequence of claim 28, wherein the nucleic acid is cDNA.
1		30.	The polypeptide encoded by the isolated nucleic acid sequence of claim 28.
1		31.	A vector comprising the nucleic acid sequence of claim 28.
1		32.	A host cell stably transfected with the nucleic acid sequence of claim 28.
1		33.	An antibody that is specifically immunoreactive with the polypeptide of claim 30.
1 2	BTF3.	34.	An isolated nucleic acid sequence comprising a sequence substantially identical to
1		35.	The isolated nucleic acid sequence of claim 34, wherein the nucleic acid is cDNA.
1		36.	The polypeptide encoded by the isolated nucleic acid sequence of claim 34.
1		37.	A vector comprising the nucleic acid sequence of claim 34.
1		38.	A host cell stably transfected with the nucleic acid sequence of claim 34.
1		39.	An antibody that is specifically immunoreactive with the polypeptide of claim 36.

1 2	BTF4.	40.	An isolated nucleic acid sequence comprising a silquence substantially identical to
1		41.	The isolated nucleic acid sequence of claim 40, wherein the nucleic acid is cDNA.
1		42.	The polypeptide encoded by the isolated nucleic acid sequence of claim 40.
1		43.	A vector comprising the nucleic acid sequence of claim 40.
1		44.	A host cell stably transfected with the nucleic acid sequence of claim 40.
1	,	45.	An antibody that is specifically immunoreactive with the polypeptide of claim 42.
1 2	BTF5.	46.	An isolated nucleic acid sequence comprising a sequence substantially identical to
1		47.	The isolated nucleic acid sequence of claim 46, wherein the nucleic acid is cDNA.
1		48.	The polypeptide encoded by the isolated nucleic acid sequence of claim 46.
1 .		49.	A vector comprising the nucleic acid sequence of claim 46.
1		50.	A host cell stably transfected with the nucleic acid sequence of claim 46.
1		51.	An antibody that is specifically immunoreactive with the polypeptide of claim 48.
1 2	NTP-3	<b>52</b> .	An isolated nucleic acid sequence comprising a sequence substantially identical to
1	• •	53.	The isolated nucleic acid sequence of claim 52, wherein the nucleic acid is cDNA.
1		54.	The polypeptide encoded by the isolated nucleic acid sequence of claim 52.
1		55.	A vector comprising the nucleic acid sequence of claim 52.
1		56.	A host cell stably transfected with the nucleic acid sequence of claim 52.
1		57.	An antibody that is specifically immunoreactive with the polypeptide of claim 54.

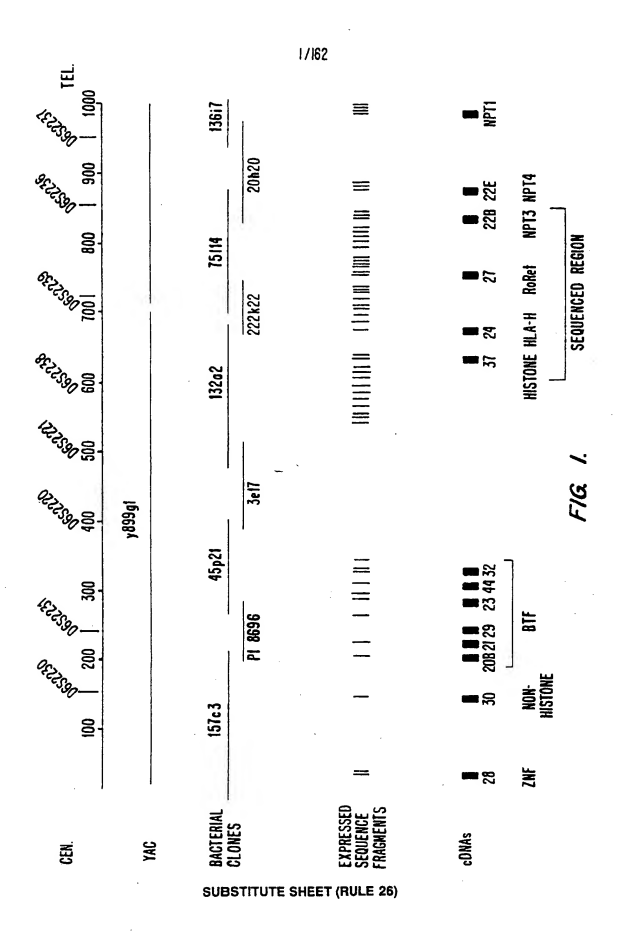
1 2	NTP-4.	58.	An isolated nucleic acid sequence comprising a sequence substantially identical to
1		<b>59</b> .	The isolated nucleic acid sequence of claim 58, wherein the nucleic acid is cDNA.
1		60.	The polypeptide encoded by the isolated nucleic acid sequence of claim 58.
1		61.	A vector comprising the nucleic acid sequence of claim 58.
1		62.	A host cell stably transfected with the nucleic acid sequence of claim 58.
1		<b>63</b> .	An antibody that is specifically immunoreactive with the polypeptide of claim 60.
1 2	RoRet	<b>64</b> .	An isolated nucleic acid sequence comprising a sequence substantially identical to
1		65.	The isolated nucleic acid sequence of claim 64, wherein the nucleic acid is cDNA.
1		66.	The polypeptide encoded by the isolated nucleic acid sequence of claim 64.
1		67.	A vector comprising the nucleic acid sequence of claim 64.
1		68.	A host cell stably transfected with the nucleic acid sequence of claim 64.
1		69.	An antibody that is specifically immunoreactive with the polypeptide of claim 66.
1 2	substa	70. antially id	An isolated nucleic acid sequence comprising at least 18 contiguous nucleotides lentical to 18 contiguous nucleotides of BTF1.
1 2	subst	71. antially ic	An isolated nucleic acid sequence comprising at least 18 contiguous nucleotides dentical to 18 contiguous nucleotides of BTF2.
1 2	subst	72. antially id	An isolated nucleic acid sequence comprising at least 18 contiguous nucleotides dentical to 18 contiguous nucleotides of BTF3.
1 2	subst	73. antially k	An isolated nucleic acid sequence comprising at least 18 contiguous nucleotides dentical to 18 contiguous nucleotides of BTF4.
1		74.	An isolated nucleic acid sequence comprising at least 18 contiguous nucleotides
2	subst	antially i	dentical to 18 contiguous nucleotides of BTF5.

1

2

	75.	An isolated nucleic acid sequ nce comprising at least 18 contiguous nucleotides
2	substantially ide	ntical to 18 contiguous nucleotides of NPT3.

- 76. An isolated nucleic acid sequence comprising at least 18 contiguous nucleotides substantially identical to 18 contiguous nucleotides of NPT4.
- 77. An isolated nucleic acid sequence comprising at least 18 contiguous nucleotides substantially identical to 18 contiguous nucleotides of RoRet.



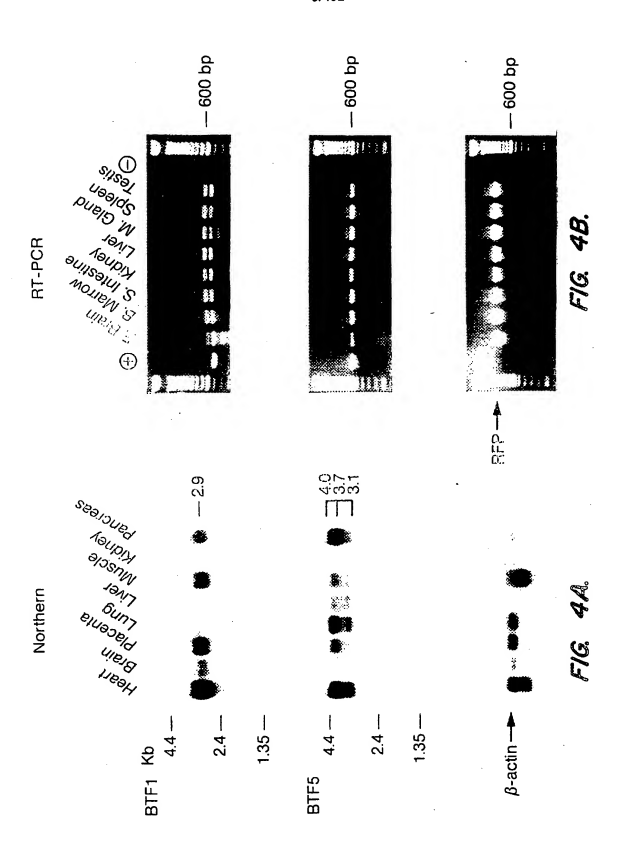
0		0000   0NA37 (		20000	25000	30000		40000 H4/g	45000	50000	55000
_!	RNA-CDI		• †	128	10.4\	Test	icular Hi		nRN	A-CDNA	<b>—</b>
137dji t	EST Pair		DE EST, y	29a03 l	<u>EST.</u> PA EST Pair					e96q01 E f6!f07 E	HLR/H ST Pair ST Pair
77602	EST Pair		' <b>8505 E</b> yd72hl2						,		
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					HL2			LH2B Psuedoge		H2a H3	H4
120000	) 12 <b>50</b> 00 • HI.I	0 13000	0 1350	00 1400	000 1450	000 150	000 155	000 160	000 165	000 1700	000 17500
H3A4	H3 Psued	do-Gene						n		IA 25/27	
			yd35d(	)5 EST p	air	yd	52a05 <sub>a</sub> r	EST	25/2		EST Pair
			•								
18000	00 18500	00 1900	00 1950	000 200	000 205	000 210		5000 220 NA-cDN/		500023	0000
					N	a/PI Tr	•				
							yr4	12a05 El	ol Pair		

FIG. 2.

BT BTF1 BTF2 BTF5 BTF3 BTF4	MAVFPSSGLPRCLLTLILLQLFKLDSAPFDVIGPPEPILAVVGEDAELPCRLSPN MESAAALHFSRPASLLLLLLSLCALVSAQFIVVGPTDPILATVGENTTLRCHLSPE MEPAAALHFSLPASLLLLLLLLLLSLCALVSAQFTVVGPANPILAMVGENTTLRCHLSPE MKMASFLAFLLLNFRVCLLLLQLLMPHSAQFSVLGPSGPILAMVGEDADLPCHLFPT MKMASSLAFLLLNFHVSLFLVQLLTPCSAQFSVLGPSGPILAMVGEDADLPCHLFPT MKMASSLAFLLLNFHVSLLLVQLLTPCSAQFSVLGPSGPILAMVGEDADLPCHLFPT  * * * * * * * * * * * * * * * * * * *
BT BTF1 BTF2 BTF5 BTF3 BTF4	ASAEHLELRWFRKKVSPAVLVHRDGREQEAEQMPEYRGRATLVQDGIAKGRVALRIRGVR KNAEDMEVRWFRSQFSPAVFVYKGGRERTEEQMEEYRGRTTFVSKDISRGSVALVIHNIT KNAEDMEVRWFRSQFSPAVFVYKGGRERTEEQMEEYRGRITFVSKDINRGSVALVIHNVT MSAETMELKWVSSSLRQVVNVYADGKEVEDRQSAPYRGRTSILRDGITAGKAALRIHNVT MSAETMELRWVSSSLRQVVNVYADGKEVEDRQSAPYRGRTSILRDGITAGKAALRIHNVT MSAETMELKWVSSSLRQVVNVYADGKEVEDRQSAPYRGRTSILRDGITAGKAALRIHNVT **
BT BTF1 BTF2 BTF5 BTF3 BTF4	VSDDGEYTCFFREDGSYEEALVHLKVAALGSDPHISMQVQENGEICLECTSVGWYPEPQV AQENGTYRCYFQEGRSYDEAILHLVVAGLGSKPLISMRGHEDGGIRLECISRGWYPKPLT AQENGIYRCYFQEGRSYDEAILRLVVAGLGSKPLIEIKAQEDGSIWLECISGGWYPEPLT ASDSGKYLCYFQDGDFYEKALVELKVAALGSDLHVDVKGYKDGGIHLECRSTGWYPQPQI ASDSGKYLCYFQDGDFYEKALVELKVAALGSDLHIEVKGYEDGGIHLECRSTGWYPQPQI ASDSGKYLCYFQDGDFYEKALVELKVAALGSNLHVEVKGYEDGGIHLECRSTGWYPQPQI
BT BTF1 BTF2 BTF5 BTF3 BTF4	QWRTSKGEKFPSTSESRNPDEEGLFTVAASVIIRDTSTKNVSCYIQNLLLGQEKKVEISI VWRDPYGGVAPALKEVSMPDADGLFMVTTAVIIRDKSVRNMSCSINNTLLGQKKESVIFI VWRDPYGEVVPALKEVSIADADGLFMVTTAVIIRDKYVRNVSCSVNNTLLGQEKETVIFI QWSNNKGENIPTVEAPVVADGVGLYAVAASVIMRGSSGEGVSCTIRSSLLGLEKTASISI KWSDTKGENIPAVEAPVVADGVGLYAVAASVIMRGSSGGGVSCIIRNSLLGLEKTASISI QWSNAKGENIPAVEAPVVADGVGLYEVAASVIMRGGSGEGVSCIIRNSLLGLEKTASISI * * * * * * * * * * * * * * * * * * *
BT BTF1 BTF2 BTF5 BTF3 BTF4	PASSLPRLTPWIVAVAVILMVLGLLTIGSIFFTWRLYNER PESFMPSVSPCAVALPIIVVILMIPIAVCIYWINKLQKEKKILSGEK PESFMPSASFWMVALAVILTASPWMVSMTVILAVFIIFMAVSICCIKKLQREKKILSGEK ADPFFRSAQRWIAALARTLPVLLLLLGGAGYFLWQQQEEKKTQFRKK ADPFFRSAQPWIAALAGTLPISLLLLAGASYFLWRQQKEKIALSRET ADPFFRSAQPWIAALAGTLPILLLLLAGASYFLWRQQKEITALSSEI
BT BTF1 BTF2 BTF5 BTF3 BTF4	PRERRNEFSSKERLLEELKWKKATLHA
BT BTF1 BTF2 BTF5 BTF3 BTF4	VDVTLDPDTAHPHLFLYEDSKSVRLEDSRQKLPEKTERFDSWPCVLGRETFTSGRVDVVLDPDTAHPDLFLSEDRRSVRRCPFRHLGESVPDNPERFDSQPCVLGRESFASGKADVVLDPDTAHPELFLSEDRRSVRRGPYRQRVPDNPERFDSQPCVLGWESFASGK KPADVILDPKTANPILLVSEDQRSVQRAKEPQDLPDNPERFNWHYCVLGCESFISGR KPADVILDPDTANAILLVSEDQRSVQRAEEPRDLPDNPERFEWRYCVLGCENFTSGR
BT BTF1 BTF2 BTF5 BTF3 BTF4	HYWEVEVGDRTDWAIGVCRENVMKK-GFDPMTPENGFWAVELY-GNGYWALTPLRTPLPL HYWEVEVENVIEWTVGVCRDSVERK-GEVLLIPQNGFWTLEMH-KGQYRAVSSPDRILPL HYWEVEVENVMVWTVGVCRHSVERK-GEVLLIPQNGFWTLEMF-GNQYRALSSPERILPL HYWEVEVGDRKEWHIGVCSKNVQRK-GWVKMTPENGFWTMGLTDGNKYRTLTEPRTNLKL HYWEVEVGDRKEWHIGVCSKNVERKKGWVKMTPENGYWTMGLTDGNKYRALTEPRTNLKL

Figure 3 (Page 1 of 2)

BT BTF1 BTF2 BTF5 BTF3	AGPPRRVGIFLDYESGDISFYNMNDGSDIYTFSNVTFSGPLRPFFCLWSSGKKPLTICPI KESLCRVGVFLDYEAGDVSFYNMRDRSHIYTCPRSAFSVPVRPFFRLGC-EDSPIFICPA KESLCRVGVFLDYEAGDVSFYNMRDRSHIYTCPRSAFTVPVRPFFRLGS-DDSPIFICPA PKPPKKVGVFLDYETGDISFYNAVDGSHIHTFLDVSFSEALYPVFRILTLEPTALSICPA PEPPRKVGIFLDYETGEISFYNATDGSHIYTFPHASFSEPLYPVFRILTLEPTALTICPI
BTF4	
BT	ADGPERVTVIANAQDLSKEIPLSPMGEESAPRDADTLHSKLIP; QPSQGAP
BTF1	LTGANGVTVPEEGLTLHRVGTHQSL
BTF2	LTGASGVMVPEEGLKLHRVGTHQSL
BTF5	
BTF3	PKEVESSPDPDLVPDHSLETPLTPGLANESGEPQAEVTSLLLPAHPGAEVSPSATTNONH
BTF4	
BT	
BTF1	****
BTF2	
BTF5	
BTF3	KLQARTEALY
BTF4	



SUBSTITUTE SHEET (RULE 26)

			CYSTEINE-RICH DOMAIN-
	52 kD RoRet	80 H	52 kD Ro MASAARLTMMWEEVTCPICLDPFVEPVSIECGHSFCQECISQVGKGGGSVCPVCRQRFLLKNLRPNRQLAMMVN RORet MASTTSTKKMWEEATCSICLSLMTNPVSINCGHSYCHLCITDFFKNPSQKQLRQETFCCPQCRAPFHMDSLRPNKQLGSLIE *** * * * * * * * * * * * * * * * * *
	52 kD Ro RoRet	Ro F	NIKKISQEAREGTQGERCAVHGERLHLFCEKDGKALCWVCAQSKKHRDHAMVPLEEAAQEYQEKLQVALGELRRKQELAEKL ALKKTDQEMSCEEHGEQFHLFCEDEGQLICWRCERAPQHKGHTTALVEDVCQGYKEKLQKAVTKLKQLEDRCTEQ
SUBSTITU	52 kD Ro RoRet	RO :	SEVEIAIKRADWKKTVETQKSRIHAEFVQQKNFLVEEEQRQLQELEKDEREQLRILGEKEAKLAQQSQALQELISELDRRCHS KLSTAMRITKWKEKVQIQRQKIRSDFKNLQCFLHEEEKSYLWRLEKEEQQTLSRLRDYEAGLGLKSNELKSHILELEKKCQG  * * * * * * * * * * * * * * * * * * *
TE SHEET (RU	52 kD RoRet	Ro Ro	52 kD Ro SALELLQEVIIVLERSESWNLKDLDITSPELRSVCHVPGLKKMLRTCAVHITLDPDTANPWLILSEDRRQVRLGDTQQ RoRet SAQKLLQNVNDTLSRSWAVKLETSEAVSLELHTMCNVSKLYFDVKKMLRSHQVSVTLDPDTAHHELILSEDRRQVTRGYTQE ** *** * * * * * * * * * * * * * * * *
ILE 26)	52 kD Ro RoRet	Ro :	SIPGNEERFDSYPMVLGAQHFHSGKHYWEVDVTGKE NQDTSSRRFTAFPCVLGCEGFTSGRRYFEVDVGEGT ** *** * ****
	52 kD Ro RoRet	Ro :	HLQVPPCQVGIFLDYEAGMVSFYNITDHGSLIYSFSECAFTGPLRPFFSPGFNDGGKNTAPLTLCPLNIGSQGSTDY HLHEQPLLVGIFLDYEAGVVSFYNG-NTGCHIFTFPKASFSDTLRPYFQVYQYSPLFLPPPGD

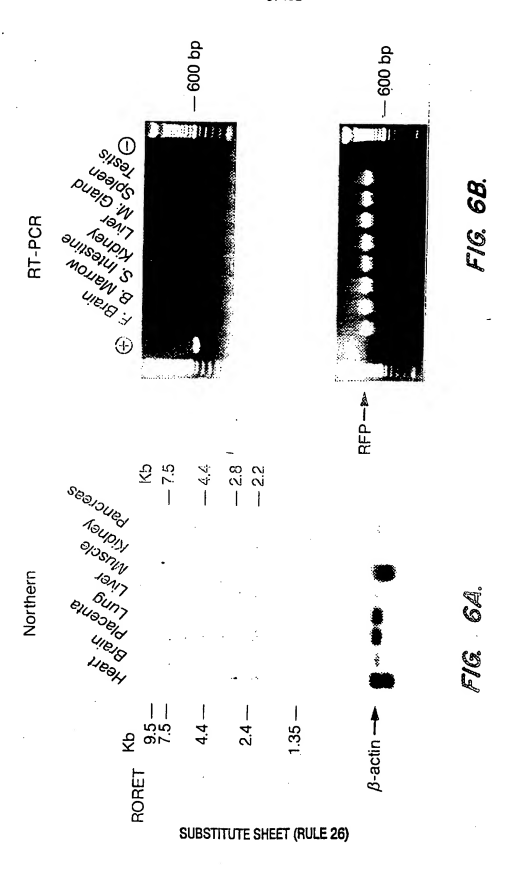
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EKEYITSSLVQQVSSSRQSLPIKAILKSLPVWAISIGSFTFFWSHNIMTLYTPMFINSMLHVNIKENGFLSSLPYLFAWICG EKEHILSSLAQQPSSPGRAVPIKAMVTCLPLWAIFLGFFSHFWLCTIILTYLPTYISTLLHVNIRDSGVLSSLPFIAAASCT EKEYIISSLKQQVGSSKQPLPIKAMLRSLPIWSICLGCFSHQWLVSTMVVYIPTYISSVYHVNIRDNGLLSALPFIVAWVIG ***	NPT1 NPT3 NPT4
TAQFEIYVKWAPPLERGRLTSMSTSGFLLGPFIVLLVTGVICESLGWPMVFYIFGACGCAVCLLWFVLFYDDPKDHPCISIS TGQFTIWAKWAPPLERSKLTTIAGSGSAFGSFIILCVGGLISQALSWPFIFYIFGSTGCVCCLLWFTVIYDDPMHHPCISVR GGQFAIWEKWGPPQERSRLCSIALSGMLLGCFTAILIGGFISETLGWPFVFYIFGGVGCVCCLLWFVVIYDDPFSYPWISTS ** * * * * * * * * * * * * * * * * * *	NPT1 NPT3 NPT4
-PMYNWSPDIQFIILSSTSYGVIIIQVPVGYFSGIYSTKKMIGFALCLSSVLSLLIPPAAGIGVAWVVVCRAVQGAAQGIVA ASVYQWSPETQGIIFSSINYGIILTLIPSGYLAGIFGAKKMLGAGLLISSLLTLFTPLAADFGVILVIMVRTVQGMAQGMAW 	NPT1 NPT3 NPT4
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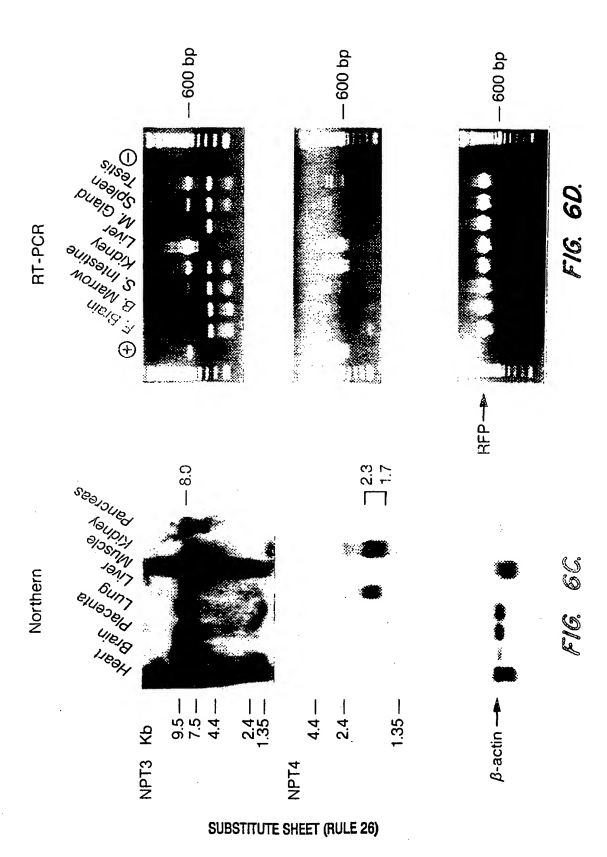
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MVGGYLADFLLTK-KFRLITVRKIATILGSLPSSALIVSLPYLNSGYITATALLTLSCGLSTLCQSGIYINVLDIAPRYSSF

NPT4

FIG. 5B.





>CDNA21 cgacccacgcgtccgaacatggcgacctaggagaaagggaagaacaattttttctcctctttttgggaagg tttgcgtctagtagtgcctgtgccctgggcagattggagagaagagggacgactggagaatcgtcgaga cactggtctcagcccagtttattgtcgtggggcccactgatcccatcttggccacggttggagaaacac tacgttacgctgccatctgtcacccgagaaaaatgctgaggacatggaggtgcggtggttccqqtctcag gaagaaccacctttgtgagcaaagacatcagcaggggcagcgtggccctggtcatacacaacatcacagc ccaggaaaacggcacctaccgctgttacttccaagaaggcaggtcctacgatgaggccatcctgcacctc gtagtggcaggactaggctctaagccctcatttcaatgaggggccatgaagacgggggcatccggctgg agtgcatatctagagggtggtacccaaagcccctcacagtgtggagggacccctacggtggggttgcgcc tgccctgaaagaggtctccatgcctgatgcagacggcctcttcatggtcaccacggctgtgatcatcaga tttttattccaqaatcctttatgcccagtqtqtctccctqtgcagtqqccctqcctatcattqtqqttat tctgatgatacccattgccgtatgcatctattggatcaacaaactccaaaaggaaaaaaagattctgtca aagaggaagaacttcaagtaaaagagaaacttcaagaagaattgcgatggagaagaacattcttacatgc tgttgatgtggtcctggatccagacaccqctcatcccgatctcttcctgtcagaggaccggagaagtgtg agaaggtgccccttcaggcacctaggggagagcgtgcctgacaacccagagagattcgacagtcagcctt qtqtcctagqccgqqagaqcttcqcttcagqqaaacattactqqqaqqtqqaqqtqqaaaacqtqattqa gtggactgtgggggtctgtagagacagtgttgagaggaaaggggaggtcctgctgattcctcagaatggc ttctggaccttggagatgcataaagggcaataccgggccgtgtcctcccctgataggattctccctttga aggagtccctttgccgggtgggcgtcttcctggactatgaagctggagatgtctccttctacaacatgag ggacagatcqcacatctacacatgtccccgttcagccttttccgtgcctgtgaggcccttcttcaggttg gggtgtgaggacagccccatcttcatctgccctgcactcacaggagccaatggggtcacggtgcctgaag agggcctgacacttcacagagtggggacccaccagagcctatagaatcaattccttggtctcacagccat gtagacaagccctggtcatctcagcagccaccgcacaacacccctggtggaagacacgccctcctcccct ctggtcacacaagagaacatcttccagctgcctctttcacacccactacagacctcagccccagttttct cctcctcactaggctgtgtttttagtagttcctttgcttgtaactatgggatgggatccaggcataggga actagttgttacacagctcccagccaagaagaaagtgtgagaagttgatgggcagcaaacctgctgttta acatcagggtgaccacattaagcccagtattccagttggcaccagaagatatggacttggaatgaggcct acagggttcaccaggatgtaagaggagaggaatccacaggaccaccagagaggagagggaaccagata ccaccataacagctaaggggacctgggagatgatggctcatttccacccagccccaggatttccagagcg cacatccacaggcctggacctgggatgaagatgaatgaagaacatggatgcacgtggatgtagtttggct caggtgtccctgcagttggcaaggagtcagtactcagtccctgagtgtggctgaaatttgaggtcctggc tgagccaaggagtaatggaccagatctacctcagtattcaagttcagtggggacaccagtggcttcaaac ttcctggtttcatgatatcttgagacgccttacaaatgatggaggattccaaagagtttttgtttatttg ggttaatatttgttggtatttatggcatttgagattgaaactaagaaatgttttaatttattacctttac ttgtgaagtgtgaaaaaaaaaaaaaaaaaa

#### >CDNA29

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181	CAGAAGAAAA	ATCAAACAGG	CATTTCAGAG	ATTGAGGCCA	AGAAGTTAA	TGTCTTAAAT
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361	AAGGAGTATG	GAGAATGAAT	TATTGCATGT	ATTGAATATG	TAGGTGACGT	GACTCACAGA
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541	GAGCACGTTT	ATTCCTGTGT	CTTGGAAGTG	TTTAGGGTGA	AAGACCTATT	' AGAGTTCTAA
601	ATGGAGATGT	CAAGTGAAAA	TGTGGCTACA	CACATTTGCA	TTTCAGAAAA	AAGGTCAGGC
661	TGGAGATGTA	AAATTGGAAG	TTTACTGCAT	ATAGATAGTC	TTTGGAACCG	TAGTATTGAT
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901	AATGGATATG	GTTATCTGCC	TGGTGTCTGT	GAAATAATTT	AAGCCAGGAA	GAGATCCTCA
961	CCAGAAACTG	ACTATGCTGG	CAACTTGGAT	CTTAGATTTC	CAGCCTGCAG	AATTGTTAGA
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1081	GACTAAGTTT	TGGTACCCAG	GCGTGGGATG	CTGCAACAAC	AAATACCTAA	ACATGGGGAA
1141	GTGGCTTTGG	AAATTGGTGA	TGGGTAAAGG	CTGGAAGAGT	TTGAGGTTCA	TACTAGAAAA
1201	AGCCAATTGT	GAAGGGACTA	TTGAAAGAAA	TATGGACATT	AAAGGCAATT	CTGGCAAAGG
1261	CTCAGAAAGG	AAGAGAGCTG	GACAGAAAGC	TTCCATTTTC	ATAGAAACTT	AGATTTATAA
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1381	GGCTCACGCC	TGTAATCTCA	GCACTTTGGG	AGGCTGAGGG	CACAGATCAC	GAGGTCGGGA
1441	GTTTGAGACC	AGCCTGGCCA	ATATGGCGAA	ACCCTGTCTC	TACTAAAAAT	ACAAAAATTA
1501	GCTGGGCATG	GTGATGTGCT	TCTGTGGTCC	CAGCTACTCG	GGAGGCTGAG	GCTGAAGAAT
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1621	TGGGATACAG	AGCAGGACTC	CACTCCCCC	GCCACACACA	CACAAAAAAT	ATATATATAT
1681	GGACATTAAA	GTCAACTCTT	GTGAGGTCTC	AGATGAAAAT	GAGGGACAGG	TTATTGGAAA
1741	CTGTAGAAAT	CACTGTTCTT	GTTACAATGT	GTCAAGAACT	TGGCTGAATT	ACGCTGTAGT
1801	GTTTACTGGA	<b>AAGAACTTAT</b>	AAGCAGTAAA	ACTGGATATT	TACCAGAAGA	GATGTCTAAG
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1921	AGAGCCGAAA	TAAAGAAGGA	ATTTTTAAGC	AAAACACAAT	CAGAACTTGG	AGATTTGGGA
1981	TAGATTTCTC	AATCTATATT	GTAAAAATTG	AGAAAGTTTT	TCTTGAAGAG	GTATGGTTGA
2041	ACAATGTTTT	CTTTTTCTTT	TTTTTTCTTG	GTTTTATTTT	TATTTTTATG	TTTTTTGAGA
2101	CAGGGTCTGG	CTATGTCATC	CAGGCTGGAG	TGCAGTGGCA	CAATCTCAGT	TCAGTGCAAC
2161	CTTTGCCTTC	AGGCTCAAGC	AATCCTCCCA	CCTCAGCCTC	CTAAGTAGCT	GGGACTACAT
2221	GTATGCACCA	CCACACCCTG	GCTAATTTTT	TGTTGTTGTT	TATAGAGATG	GGGTTTTGAC
2281	ATGTTGCCTA	GGCTGGTCTC	TAACTCCTGA	GCTCAAGTGA	TCTGCCCTCC	TCAGTCTCCC
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3001	AGAGTACAGA	TGGGATAGGG	TGGGGGTGGG	AACATGTAGT	CAAGGCTGAC	TCTACCTGTT
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2247						
3241	TTTCTTAAGA	CCTAACAGAA	TTTGCCTTGC	CAGGTTTTGG	ACTTGATTAG	GACACATTAC
3301	ACCTTCCTTC	TTTCCTATTT	CTCCATTTTC	TAATGGGAAT	GTCTATTATG	CCTGTTTCAC
3361	CATTGTACCT	TAGAAGCATG	TAACATTTCT	GGTTTCACAC	GTTCAAAGCT	GGAAAGGAAT
3421	TTTGTCTCTG	GATGAATCAC	ACATTGAGCC	TCACCCGTAA	CCTGATTTAG	ATGATTTTT
3481	AGATGACACT	TTGAACTTTA	GAATTGATGC	TAGAATGAGT	TAAGACTTTC	AGGGGGCTGT
3541	TGGGATGGAA	TAATTTTTT	TTTTTTTTTG	AGACGGAGTC	TAGCTCTGTC	GCCCAGGCTG
3601	GAGTGCAGTG	GCACCATCTT	GGCTCACTGC	AAGCTCTGCC	TCCCGGGTTT	ATGCCATTCT
3661	CATGTCTCAG	CCTCCAGAGT	AGCTGGGACT	ACAGGCGCCC	GCCACCACGC	CTGGCTAATT
3721	TTTTTTTTAT	TTTAGTAGAG	ATGGGGTTTC	ACCGTGTTAG	CCAGAATGGT	CTCGATCTCT
3781	TGACCTTCTG	ATCCGCCTGC	CTTGGCTTCC	CAAAGTGCTG	GGATTACACG	TGTGAGCCAC
3841	CATGCCCGGC	TGGGATGGAA	TAAATTTATC	TTGTATGGGA	GAAGGACATA	CATTTTGGCA
3901	GGTCAAGGAC	AGAATGTTAT	GGACTAAACT	GTGTCCCCCA	AAATTCATTT	ATTAAAACCC
3961	TAAACCCCAG	TGTGACTGCA	TTTGGACATA	GAGCCTTTAG	GGGGTACATA	AAACTAAAGA
4021	TCACAGGATA	GGGCCCTAAT	CCCATTGGGG	CTGGTGTCCT	TACAGAAGAT	GAGACACTTA
4081	GAGCTCTCTC	TCCACGCAGG	CACCAAGGAA	ACACCATACA	AACACACAGT	GAGATGGCAG
4141	CCATCTGTTA	GCCAGGAACA	GATTCTCACC	ATAAACTATG	TTGGCACCTT	GATCTTAAAC
4201	TTCCAGGCTC	CAAAACTGTG	AGAAAATGAA	TTTCTGTTCC	AAGCCTCTTA	GATATGGAAA
4261	AAAAGATTCT	GTTGTTTAAG	CCATCCAGTC	TCTGGTATTT	TGTTATGGCA	GCCTGAGTAG
4321	GCTAAGACAA	TGAAGGATGT	GGTAAAACTT	TACGTCCCAA	CCACATACCA	AAGAGGCTCC
4381	AATTTAGCAT	GCTTTCTTCT	TTCAACTGTA	GGCAATGTGC	ACAAGTTCTA	AATCCTAAGA
4441	CATGTTGGCT	CCTTTACTCT	GCCCAAACTA	CAACTCAAAC	AAACAACTGT	אמיית מיית מ
4501	CATCCAATGA	AGTTCTGACA	TTTCTTCAAC	ATGAGTACAG	TAATTCAATG	CCDCDCDATT
4561	CATTTTATTT	TGAAATCTAC	ATGCCATATT	CCAATTTCTG	TTGAAGATGC	AATCCTTATA
4621	TTTATTCTTT	TTAATATAGA	TTTATCAGAC	TEGECECECT	GGCTCATACC	TOTARTOCTA
4681	GCATTTGAGA	GGCTGAGGTG	GGCATATCAC	CTGAGGTCAG	GAGTTTGAGA	CCACCCTCCC
4741	CAACATGGTG	AAACCCTGTC	TCTACTATAA	דממממדמדמ דממממדמדמ	TAGCTGGGTG	TCCTCCTCCT
4801	TGCCTGTAGT	CCCAGTTACT	AGGGAGGCTG	ACCUPACIA ATT	GCTTGAACCT	CCCACCACCA
4861	GGTTGCAATG	AGTGGAAATC	GCACCAGTAC	ACTCCAGCCT	GGATGACAGA	CCAARAMAR
4921	AAATAAATAC	ATAAAATAGA	TTTATCACTT	TATCANTANT	ATAGTTTTCT	GCAAAATAAT
4981	TAAATATAGG	TAATGACTGT	CCTTTAGTAC	ATCARIAGI TATCARIAGI	GATGCTCCTC	TITCTAGGIG
5041	TGGTACAATA	TTAAGTATTG	מדמממדממם	CACAATCCTC	TCCCTACACA	TIACIIGGIT
5101	TTCCATTTGC	TCATCTCCAA	TATECACEGE	AAATTCTCAA	ATTGCTAATA	TGAGCACTTA
5161	ACACATGCAT	TATATTCAAC	AGGAATATAT	AAATICICAA	MIIGCIAAIA	ATCTTGTAAC
5221	ATGACAAACC	TTTAGAAGGT	TTGTDTTTN	CCTTANANTA	TARMATTIA	GGATCAACAG
5281	ATAAAATTTC	TAATACTTTC	TTOTATTTAK	CCTIAAAAIA	ARRITITITA	AAAATTGGTT
5341	GTTCAAATGA	TTTACAGAAT	TOTALARCE	TATACACAGG	AAAATATAAT	TCTTATAAAA
5401	GATATTGCTA	CATACACACA	CARAMMOIG	AATAGAGATG	ATGAATGAAT	TAAAGGAAAG
5461	AAACTGATCT	CATAGATITG	ACAMATITAMA ACAMACCOMPA	AAGGGAAATT	ACGATIGITG	ATTTTGTGTT
5521	ATATCTCAGT	AAATTCCTCA	CACAAACTIM	ACTOCCAGA	AATGATTTTA	TCTCAGCCTC
5581	TGGGAGACCT	CTACCTTTAC	CATCCTCATC	AGICCCIGGI	GCCCAGGTGC	CTTTGGTAAT
5641	GGGCCATTCA	GGCAAGGGAG	ATCARACTO	CACTOGOCOC	AATTTAAATA	GTCCTCCCCA
5701	CGAAATTCAT	TGCTCDATAG	ATGAAAACII	GCTCAAGAGT	TGGAATCCAA	CTGAAGCTAC
5761	AGTGGGCATT	TCAAAGTAGA	ATAMITITUC	CTGGAAGTAA	CTAGGGCTTT	TGAATATAAT
5821	CGAGGAATGT	CCAMAGIAGA	AGGIAAAGIA	TTTTGGAGAT	GAGGAGACAG	GACAGAGCTA
5881	CGAGGAATGT	CCTTIGCTIA	THOMOTOR	TCTTAGCAGT	ACCTCTTAGG	TAAGAACTGG
5941	TTAACTGGCA	ACCUSACAAC	TOTOTGAAG	CTCCCTTTGC	TTAGGGACTA	GGCTCTTAGC
6001	AGTACCTCTT	WARTWARE .	CAMPAGE	ACACCTTCTA	TGTGTCTGAA	GCTCCCAGAA
6061	CAAACTGCCA	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	ACTORITY	ATATAGTTTC	TITTTTCTTG	TTACTTTTTG
6121	TTTTGTTGTT	TITITIGAG	AGTOTCACTO	TCACTGCAAC	CTCCCCCTCC	TATATTCAAG
6181	TGATTCTCTT	CUNTUMBER	CCCGAGTAGC	TGGGACTACA	GGCGTGCACT	AGCATGCCCA
6241	GCTAATTTTT	BCCCTCCTCTA	GIAGAGATGG	GGTTGGTTTT	TTTTTGAGAC	GGAGTTTCAC
6301	TTTGTCGCCC .	AGGC TGGAGT	GCAGTGGCAC	GATCTTGGCT	CACTACAACC	TCCACCTCCC
6361	GGGGTTCAAG	CCCACACAC	GCCTCAGTCT	CCTGAGTAGC	TGGGACTACA	GGCGCCTACA
	GGTGAACACC	GCCACACCTG	ACTAATTTGT	GTAGTTTTAT	TAGAGATGGG	GTTTCGCCAT
6421	GTTGGCCAGG	CIGGICICAA	ACTCCTGACC	TCAGGTGATC	TACCCACCTC .	AGCCTCCCCA

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6481	AGTGCTGGG#	TTACAGATG	T GAGACACCAG	ATCAGCCTC	ם הממה המידיים	TCTATTGGAA
6541	AGAGAAAACA	CTATTAGCA	A CCTATTAGTO	מיייימית מתמת בי	TACTTARTO	CTTCCTTAGT
6601	AATAAACCAA	CTCTCTACAL	A CAAAGTGCTT	CCTGGCTGC	TACTIMATES	ATTCATTCAG
6661	TTCAACATTI	TCTCAATGC	CAACAGCCAA	GTGTCTCTT	TATCCCARC	TCTATGCTGA
6721	TTATCAGTAT	TTGAATAAG	A GGGGGTCTAC	ATCTTAAGT	CTCCTTNACA	TGAAAGCCTC
6781	TAGGTTAACA	AACTTAACA	AATGTATCAT	י דרברדבתפון דרברדברדב	TOCITARGA	TACAAAATCT
6841	TGTTATTGGA	GCCCAGAGAG	AAGAATTGAA	ATTCAACTAC	TOTOTOTOTO	CTTTTCTCAC
6901	TCACCACAAT	AAGTCAGTTC	CACCAAGTCT	TGTAGCTCTT	TOTOTOTOTO	TGTTTTCACG
6961	TGTCCCTTTG	TTTTATTTG	CACACCCTAR	ATABABATTO	TACTGAGCCA	TTTTCCCTGG
7021	GTTTACAGTA	TTAATACATT	GTCAAGATTT	, שכניהניהיהנים היידיה	CTACIGGCIII	TTTTCCCTGG
7081	TACCTTTCCT	CCTTCCCTT	AATTCTTCAG	AGGTTAGAA	GLAGATICCC	ACATTCTGGT
7141	ATGTGGACAA	AGTTTACCCA	TTATGTATGG	ATGTTTTACT	. GCCWIIWGIW	TTCTGACAAT
7201	AATCTCTTAA	GGAGGTGTGG	TTATAGAATA	GTCAGCTGTT	. CITICIMITI	TTTTCCTGGC
7261	CTTACAACTT	AAGTTCTTTA	AGCTGTTTCT	TAGTTTGCTC	DIDMIDMALA	TCGGAATAAG
7321	GATAAAACCT	ATCTCTTAGA	TTGTTGGATT	משייה מבשייה ממ	AICICAAAAI	AGCTCATGAA
7381	ATGTGCCTGG	CACACAGTAG	TGCCTAATAA	ACCATCTCTC	TTATTCAGCC	AGCTCATGAA
7441	TTTCAGAATC	TACACTTGCT	GAGCCAGGTT	MCCAICICIC	CAAGGTGAGC	TGTTTTCTGA
7501	AAGGAAGAGA	TGGAGGTAGG	AAGAGATTAA	GCCCTAGCCC	AAGGTCACAC	AAAAGCATAC
7561	AGCTGGAATC	AAAGGCAATT	TGGTCAGTGA	ATARARACCA	TTCCAAGGCC	ACCGATTGGG
7621	TTCTAACCTT	AGGATCGAAA	TTCTCGGACA	TACACCARAT	GCTGGGGGGG	CATAAGGCAA
7681	TCTTCTCAGC	CCAAGAGCCA	TGTGAAACCA	GACCTTCAAA	TCTGATGATT	GAAAATCCGG
7741	CTGCCCATTA	GAATCGTTGT	AATTTAAAAA	TACCCTCGGA	AAATTCTAAT	ATICTICACIONA
7801	CAAAGGTGAT	CATTTGCTTT	TATGCCACTT	TGTTTTCACC	CAAATGGGAC	AIGIGGCTAT
7861	TTTCCTTTGA	GAGTAGTTGT	AGGGAAAGGA	GGGGGTGGAG	GGAGGGAAGA	ATCCAACCCT
7921	CTGGATCCGC	CCTGAGCCGG	TGTCAGTATC	TGGGAAGTGG	GAGGCGCGTC	CCGGAAAAGG
7981	AGCTTCTGCT	AGGATTATTA	TCTCCTGCCA	CACACTCGGA	TTTGAAGGCT	AGCAGTAAAC
8041	CAATGCAAAA	CGCTTCAGTG	GAGTTCCAGA	AGCGTTAGAC	TAAACGACTG	CCHAACGAAA
8101	GCCAGTCTGA	GCAGCTGGGC	GCAGATGCAT	AGGCAAGACT	TAGCCCGCCT	ACACOMOROCO
8161	GCCCACTTAA	TTCCGATCAA	AGCAGAAACC	GGCCGGGCGC	GGTGGCTCAC	CCCCCTATC
8221	CCAGCACTTT	GGTAGGCAGA	GGCTGGCGGA	TCACCTGAGG	TCAGGAGTTC	GACACCACCC
8281	CGGCTAACCT	GGTGAAACTC	CGTTTCTACT	GGTGGCGGGC	GCTTGTAATC	CCATCTACTA
8341	GGGAGGCTGA	GGCCGGAGAG	TCGTCTGAAC	CCGGGAGGCG	GAGTTTGTAT	GCAGTGAGCC
8401	GAGATCGCGC	CACTGCATTC	CAGCTTGGGC	AACAGGAGCA	AAACTCCGTT	TCANANACC
8461	AAGCAAACAA	ACAAAAAAAT	GCAGAAACCG	AGATCCGGAA	GAAAACCTCG	GCGAGATTCA
8521	CAGAATCCAG	GAAAATAGGT	CTCTAGAAAT	TTGTCCATGG	TCCCAGATCT	CCATTTCTTC
8581	TGGGTGGGGC	AGCTGTTACC	AGATCCCTAG	AAGCAAAGGT	TTTTTTGGGG	GACCGTGTCT
8641	CACTGTTGCC	CAGGCTGGAG	GGCAGTGGCA	CGATCTCGGC	TTACTACAAC	CTCCCCCTCC
8701	CAGGCTCAAG	CGACTCTCCT	GCGTCAGCTT	CAAGAGTAGC	TGGGATTACA	AGGTATGTGC
8761	CACCACGCCC	AACTTATTTT	TTTATTTATT	ATTTTTTTTT	AGTAGAGAGG	TGTTTCACCA
8821	TGTTGGCCAG	GTTAGTGTCG	AAGTCGTGAC	CTCAGGTGAT	CAGCCCCCTC	GGCCTCCCAA
8881	AGIGGTAGGA	TTAGAGGGGT	GAGCAGAAAG	CAAAGGTTTTT	TCACTCCCCA	Checocoona
8941	TUTATTTCCT	TTTCTGCCTG	TAATGGCAAC	CTAGACGCTT	GAGCTTCTTA	AAATACAACA
9001	GIMMGIIGCM	IGICAGGCAC	CGTTCTACAT	TAGGGACATT	AGTCTGTTTT	ACAGACACCT
9061	TICAACICCC	TGGTTAACTT	TTAGGTAATA	TACTCTGCAC	TTTAGCAGGA	ATCCCACCTA
9121	TAACTCTCAC	AGAATTAGGA	AAGTGAGGCT	GCCTACAGCC	TAAATTCACA	2222222222
9181	CGGGGGACTA	GTCGGAGGAC	CAAACAAGGT	TACCAACACG	TTAGAGTTTT	CCCTTCDDTT
9241	IACATITITA	AAGTAATCAC	AACGAAGTGT	TTAGATCACG	AGGCATCCCT	CCATCTAAAC
9301	IGITAGGCAC	TAACTATGGT	CGATCTTACA	AAGCATTAAC	TAGAATATTT	CTTTAGAGTA
9361	IGATAGTACG	TAACTGACCT	ACTATTACAT	ACAAACAGAC	CAACCTTTAG	TAACAGCGCT
9421	CCCCAAAAAC	CGAAAAGCAG	TAATACGCTT	TGCTCAAGGT	TGGCATAAA	TTDACTTACC
9481	TIAGTGCCTT	TTTTCCTTCT	ACCTACAAGC	AGTGAGGTTA	GCTCTTCCTT	TCDDDCCCTD
9541	GGGGGCTCT	GAAAAGAGCC	TTTGGGTTTG	ATAGCGTTTC	CGGGAGCTCA	CATACCTCTC
9601 9661	AAATCACTTG	CCCLIGGCCT	TGTGGTGACT	CTCGGTCTTC	TTAGGCAGAA	GCACGCCTG
700I	GATGTTAGGA	AGGACGCCGC	CCTGAGCAAT	GGTCACCCGG	CCTAGCAGTT	TGTTGAGCTC

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9721	CTCGTCGTTG	CGGATGGCCA	GCTGCAAGTG	GCGCGGGATG	ATGCGAGTCT	TCTTGTTGTC
9781	GCGAGCCGCG	TTGCCGGCCA	GCTCCAGGAT	CTCGGCGGTC	AGGTACTCTA	ACACCGCCGC
9841	CAGGTACACC	GGCGCGCCTG	CCCCAACCCG	CTCTGCGTAG	TTGCCTTTAC	GGAGCAGGCG
9901	GTGCACTCGG	CCCACCGGGA	ACTGGAGACC	AGCGCGAGAA	GAGCGGGATT	TCGCTTTGGC
9961	GCGAGCTTTG	CCTCCTTGCT	TACCACGTCC	AGACATTGCA	ATCAGACAAA	AATCACCAAA
10021	ACCAGCGGCC	TAAGCTCACG	AGAAAACAAA	CAAAATCAAG	AAATATGTAA	AACATGGCCG
10081	CTTTTATAGG	TAGTTCCTGG	GGAGTAAATC	CGACTTTTTG	ATTGGTCGGT	AGCAAATGCT
10141	AGTCAGATAG	CCAATAGAAA	AGCTGTACTT	TCATACCTCA	TTTGCATAGC	TCTGCCCACG
10201	GATGACAACT	GTGCAGTTTG	TCTTCCAATT	AACTAAGAGG	TACTCTCCAT	CCCTCATTAG
10261	CATAAAAGCC	CTATAAGTAG	CAGAAATCCG	CTCTTTACTT	TCGACACATT	TCTGGTGTTT
10321	TAAGATGCCT	GAGCCAGCCA	AGTCTGCTCC	CGCCCGAAG	AAGGGCTCCA	AGAAGGCAGT
10381	GACCAAAGCG	CAGAAGAAAG	ATGGCAAGAA	GCGCAAGCGC	AGCCGCAAGG	AGAGTTACTC
10441	TGTGTACGTG	TACAAGGTGC	TGAAACAGGT	CCATCCCGAC	ACTGGCATCT	CTTCCAAGGC
10501	CATGGGCATC	ATGAATTCTT	TCGTTAACGA	CATATTTGAG	CGCATCGCGG	GCGAGGCTTC
10561	CCGCCTGGCG	CATTACAACA	AGCGCTCGAC	CATCACCTCC	AGGGAGATCC	AGACGGCCGT
10621	GCGCCTGCTG	CTTCCCGGAG	AGCTGGCCAA	GCACGCCGTG	TCGGAGGGCA	CCAAGGCCGT
10681	CACCAAGTAC	ACCAGCTCCA	AGTAAACATT	CCAAGTAAGC	GTCTTAACAC	CTAACCCCAA
10741	AGGCTCTTTT	AAGAGCCACC	CAGATACCCA	CTAAAAGAGC	TGTGGCCAGA	CGCCAAATTT
10801	TATTTGGCGG	CGGAGGGGTA	TTAGAATATA	GGAACTGGAG	AGGGGTGGGG	ACAAGTGTTG
10861	CAGCTTAGAG	AGGGACAAAG	GGTCCTGAAC	CCGAAAGAAG	CCAGCCATTA	AAAATGGCTT
10921	TGGGGTCAAT	TCGTTGTGCT	TAAATTTAAA	ATGGAGACAA	GCGGCCATTT	TGCTAACTCG
10981	GCGTTCCCGG	AAGAAACCGC	AGGCTCGCTT	AGGTTTCAGA	CCCAGCTGTC	TGTCCCTGTC
11041	TACGTCGCCA	GGATCAACGG	TTGCCGTAAT	GTCATAATTT	CGCCACCAGC	TTCTAGCCAA
11101	TAGGCTGTCC	TGTCATTTTA	AATATTAACC	AATCGAGGGA	AAGCTGTTTT	GAGACTCTGA
11161	TTTACATAGC	GGACCGGAGT	GGGAACCTGG	GCAGTAACTG	CCTAAGGAAG	GACTCCCCCT
11221	CTGTTTTCGT	GGCGCACACC	TTCGTAGTAT	ACTGAAGGGT	GTGTCTCCTG	GGTTTCCAAC
11281 .	TGCCCCGGTA	ATAGTCTTTT	AACCTAATAT	GCGTCAGTTT	TGATAACAAC	ACTANGGCAG
11341	TACAGAACTA	AAGATGTAAG	CACTGCGCCA	GATGTTGCTT	CATACATCTT	ATTOTOTO
11401	ACTGGTTTAT	TCAAGATTCA	AATCAAATCA	AATTTTGCTT	GAATCCCAGT	GCTCAGTCAG
11461	CCATAAATGG	TGTGTTGCCT	GATTGAAACT	TAAAATCTCC	GTAGGGGGCT	TGTAACATGC
11521	AGACAAGTTT	GAAAGTTGCT	TTAGGAGAAG	CCAACTCTTA	ACTGCTGGGT	AAATTCACAA
11581	GCCTTCGAAC	ACTGAACTGA	AGGCCAGTAA	GGACTAGGCG	CTGGGTGGG	CACAATCAAC
11641	AGGAGACGTC	ATTAAACTTA	GCACATACAC	TGTATCTCCT	AGAGGACTCT	CCCTTCCTAC
11701	ACAACTGCAG	GCCGCTTTGT	GGCCTGGGAA	ATTCCACATT	CCCTTAAGTA	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT
11761	GGTCTTTTCC	AGGTAAAGAT	TTTAAGATGA	AGGGTTAGAC	GTAGTCTACC	TITIACICAL
11821	TTCAAGTCTA	GAACACGTTT	TTAGCACCTA	GAAGTTTGCT	TTCTCCATTA	AAAACCCCCA
11881	ATATACAATA	AATAAAATTA	GTGTTAAAGC	AGATTTTTAC	AAACTTAAAT	ACCATCTAAT
11941	TTAGGTTACA	GTTATTTAAC	ATAAGGACTG	TGTGATCTTA	AATCTGCAAT	TTCTTTCACA
12001	CCTGGGAAAT	AAACTAAGGC	CTGTCTTTGG	TGCCAGACAA	GGCCTTATAC	TTCDACACTC
12061	CTGTGCAATC	ACAGGCTGCC	TTGCCTAGAT	AACTTATCTG	AGAAATTCTG	ATCACACACTG
12121	AAATTTCCAG	AGTCCCTCAC	AAGTAAATTT	Julahalahin (adalah kata kata kata kata kata kata kata ka	TTTTTTTTT	TIGAGAAAIG
12181	GAAGTTTCTC	TCTTGTTTCC	CAGGCTGGAG	TGCAATGGCG	CGATCTTGGC	TCACACGAC
12241	CTCCGCCTCC	CGGGTTCAAG	CCATTCTCCT	GCCTCAGCCT	CCGGAGTAGC	TCACAGCAAC
12301	GGCATGCGCC	ACGACACCCT	GGCTAATTTT	GTATTTTTAG	TAGAGACGAG	COUGATIACA
12361	GTCGGTCAGG	CTGGTCTCGA	ACTCCGGACA	TCAGGTGATC	TGCCCGCCTT	CCCCCCCA
12421	AGTCCTGGAT	TACAGGCTTG	AGCCACCGCG	CCGCCCTAA	ATGGTTTTTT	OGCC1CCAA
12481	GCCTCTAATG	GACCTGGTCA	CTTATTCCCA	TTCAGACTGA	CCGCTCTCCT	ACCTCCCAAC
12541	TAACTAATCA	GTGTAACCAA	AATCTGCAAA	CAAAATTCAG	TATTCTTTCC	CCCCCAGC
12601	CCCTTTCTCT	TACATAGATT	ATGTTTTTGC	CTGTGTTAGA	TGAAATAATT	COGCCITIC
12661	TTCTCTCTTC	TGTACAAGTA	CCCAGTAAGC	ΑΔΑΤΤΣΤΙΛΟΛ	CTTCTTGGTC	Z-MATTACTIC
12721	GAATTTTCCA	CCAAGACAGT	GTTTATGTGA	GTCATACAAT	PUCTORIC	WILL TAIL TO
12781	GTCTTGGAAA	CAGGTTGTCT	ATCCCTGGAC	CCTTTCACTT	TTCTCTTCA	TOWARTO TO I
12841	CTTTTGCATG	CTAAAAGTTT	ATCGTCCGCG	TTTGAGII	TICIGITOR	TAATTCCIIIGG
12901	TGGCTGATTG	GTTGCATATT	GGTGGCAGTA	GTAGAATTTC	77dd11W11C	TARIIGGACI
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12961	ATCATTAAGT	GATTAGTCAG	TGGAGAGGAC	AGGAAATCTG	GTTTATTTAT	TAACCTTTTT
13021	TTGGGGTGTT	TTTGTTTGAA	GATGTTGATA	TTCTCTGTGA	GGACACAGGG	TTAGAGTTCC
13081	TGTTTTTCTT	TCTGACTTTA	CATGGGATTT	GATGTTTTGT	GCTTGTATGC	CTCTTTCCAC
13141	CTTCCAAAAC	TTGTCTTTTT	'TGAGTCCAAA	TAGTTGTCGA	TATCTGCAAA	ACCAGTATTC
13201	CTGTGTTAAG	ATGATATGAA	TATAAAATGG	CTGCCCTGTT	ATAACTTTTG	АСТТТАВСВВ
13261	AGTGTTAGGA	CTAACAGGAG	ACAAAAAGGA	AATCAAGGAA	ACCGAATGTC	TGGTCTCAAT
13321	AACTGCTATG	GCAGAGGCTC	TACAGCTTAT	TATTAATTTT	AGTAATTTCA	CATTATTCCC
13381	CCTTCACGTT	CTTTAAGTAA	GGTTAGAGGA	CAGAAGAAAC	ATAATGTTGT	TACAAATTCC
13441	ACTATTGAGT	CAGGGAAAAA	AAAGAGTGCT	TTCAATATCT	GAATAAAACA	AACATTTAAT
13501	ATTTTCTAAA	CCTTAACGAG	TTTATTGTAA	GGGATGTGAT	GCTGGAAACT	ACCHARCTAC
13561	AATTTTCTTC	TAAACTGAGA	ATCAGAATTA	TTCATATTCT	CAGCAGTGGT	CCCACCTCAC
13621	GGACTTCTGA	TCTTAATTAC	ATACTTTTAT	TTCTTTAACT	GATCAACATC	CTARATACAM
13681	AACCTATGGC	TCTGTTTTTA	CCCACTTTAA	ATTCTGTTCT	ATTACCACCG	TTNAMIAGAI
13741	TAATTGGCAA	TAAGATTGAG	ACTATCTTTT	delinialabelahahahahahahahahahahahahahahahahahahah	GAGACAGAAT	TINGCITICC
13801	GGCCCAGGCT	GGGGTGCAGT	GGCACAATCT	CGGCTCACTG	CARCHORAL	TITIGGTGTGT
13861	CTAGCAATTT	TCCTGCCTCA	GCCTCCCCAG	TACCTCCCAT	TACACCTCIGC	CICCAGGII
13921	TGGCTAATTT	GTGCATTTTT	AGTAGAGATG	GGGTTTCGCC	ATCTTCCCCA	A A COMPANIE
13981	GAACTCAGGT	GATCCACCTC	GGCCTCCCAA	AGTGATGAGA	TTACACCCC	AACIGGICIC
14041	GCCCAGAAAA	GACTATCTTA	TTTTATGAAT	TTAAATTAATT	CTCAAATTAM	GAGCCACCGT
14101	GAATTAATAA	ΑΤΤΑΤΑΑΤΟΤ	AATCTTAAAT	TIMESTACI	GIGAAAITAT	CCACTTAAGG
14161	CATCAATTTA	AATAAAAACT	CATTTGTCTA	111AG11GGC	TIACATAAAG	ACTTAAAATA
14221	AAATGTGCTA	CCTCTTTAAG	TTCTAATTAA	CACAAAAAA	COMMITTE	CTTGTGCTTT
14281	AGTGGTCTTA	GTTABCAGCT	TAAAGTATTT	TOTARARA	GTTTAACTGT	GAGTTTCATT
14341	AACTTAAAAA	TATTAATACC	TCTTTTATTA	COMMINMEN	ATACTTCACA	ATTTTTAAAT
14401	ATCTAATCAA	Cydulalalalata	GGACAAATTG	COTTANTANT	MIAAGGAAAA	TATATAATAC
14461	מייים חיים מייים	TACTCTAAAA	ATAATATTAG	GCTTAATAAT	TTCATTTTAA	AAATGGCTTC
14521	TTCATATTCT	ממחמממממ	AACAAAAGCT	CAGAATATTA	TAGTATACAC	AAGTTTAGGG
14581	ACTACTTCTA	CTCCTTACAT	GAGTTAACAT	AATTTAACTT	GCATTTACTA	AATTTCTTCC
1:4641	TATTCATTCA	ACCANATORA	BEGREEAT	CACTTTATTT	ATTATTCTAA	AATTGTAAAT
14701	TATICATION	TO ATTACK	ATGATAATAG	ATAATGTCAT	TTTTAAAAAT	GGAATTAAAT
14761	CATARCTTER	ACARTMIAAG	GATTCAATGT	GTGAGCTTAA	GTACTGAGTT	CACAGTGTAT
14821	GATACCTIA	CARTTERES	TGAATATTAT	TAAATTGAGT	AAATTAATTC	TCAATCTTTG
14881	TTTTATCCAA	ATABCATTAM	ATTGGAGGGT	ACAAAATACA	AATCACAAGA	AACAGTGTAG
14941	CATATCATTC	CCTTACAATA	TACACAGTTT	AGAATAACCA	TTGATAAACA	GATAAGAGAA
15001	ACTOTATACO	TCTCCCCCCTA	GATACTGTTG	CTTTCGCCAC	TTTAGATTTG	TAAATCACGT
15061	CATCCCTATC	CCCCAACACA	GAGGACCATG	CAGGTTTTGG	ATGACTGCCT	CTGTTTTCGT
15121	CTCTCACTCT	ATCA ACTA CT	ATTGCCTGCT	TTGTTTAAGG	GCTATGGTTA	ATCCAAACAG
15181	CTACCTTCACTCT	ATCAAGTACT	ATAGCTACAG	AGAAACACAA	GTAAGCATTC	GAGATAATGA
15241	TTTTACTAGE	CCTTTACTTA	TTTAAAAAGT	TGTTACTGTT	TGTTAATGTG	GTACATTCAA
15301	ATCATTTANA	ATTGTCACTC	TAAAATAAGA	CTTCAATCTT	TTTCTTATTT	TTATATAGCC
15361	ATGATTTATA	CATATOTT	AATGTAATAA	CCAATCTTCT	CTGACAACAT	TATAACAATG
15421	TGGATATCTC	CATTITCAGT	ACTTCAAACA	ACAAATACTG	CTTTTATACT	TCAGAGCAGA
15481	AAAAATACAC	CITCCCAGTG	TAAACACATT	TGGAATCTCA	CTGAGAAATA	CACTATCACT
15541	AAAAATACAG	TTCTGAGATT	CATTAAAAGA	CCTCCAGAAT	TCTGGAAGTA	GGAAGTTTCC
15601	TCTTCAAAGT	CTACAGAGGA	AGATGAGGTC	TGAAATAGAC	AGCTTCTTCC	TTCTTTTACC
15661	TGTGGTATTA	TICIGITITG	TCCTTTTCTC	CATTATCTGT	CTTTCCAGTG	ATGAAATTTT
15721	GATCTGGCCC	TCCCAAGTAT	TAAAAAACAA	GCAAATAAAC	AAATCTCAGT	TATATTTTAC
15781	TAAGATATTG	GCATGCTAAC	TTTTTGCAGG	TITGTAACAA	GGACCTTTAT	AACTTGACTA
15841	AAAGTTCCTA	AATAAGAATA	TTTACTAGAA	AATTTATTTC	TGCCTGTGGC	CCACATTTGA
15901	GTCAAAATAA	TCAATTAGGA	AAAATGAACT	TGTTTAACTA	AAGTTGACCA	AACTGATCTT
15961	TGACCAAACT	GATCTTTGAG	ACCTATTCAT	CTAAGACAAG	CCAATTAAAT	TCTTGGAGAC
16021	AATTTGTACT	TTAAGGAATT	CTTATAATAT	TTGTAATTAC	CCTCATAACT	TTTTTTTTTG
16021	CCCTACTTCT	GIGCTTCTCT	AATATGCAGA	TTATTAAATG	TTGTTACAAA	GCCATTGTCA
	AAAAAACAAA	AAACAAAAA	CTAAACAAAC	TCACATGGTT	AGACTTGCTC	CTTTATGAGA
16141	TATTTTTACC	AAAAATGGAG	GAGTTGAAAA	ACTCTGGTGC	CAGAAATCGT	GAAGACATGG

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16201	CCTACCTAAC	ATGGAAATGT	TGGTTGTCAG	TGGAAAATAC	TACACAGAGA	TAGCCATAGT
16261	GCTGCACAGC	CAATCTTAAG	TGTTTCTAGA	GAATCACTAA	TTGTTTCTAG	AGAATCACTA
16321	ATTGTTTTCT	TTTAACATTC	TTGGTTTATA	CAAGAAGAGA	GTATCCATAC	TAAACTCTTT
16381	TCTACTGAAA	. ATAATGTGCA	AACATAACAT	CCTATTCCTA	GACAGTTTGT	AGTTTTTTTC
16441	TCCCATTTCT	ATTTTATAAA	TCATCTTTTT	AAAATACTTT	GTTGAGTGAA	ATCAGTCCAT
16501	TGCTTGATAT	ACCTTGAGCA	CAAGTAAATA	GTATGCCAAA	AATTAAATGT	CTTTCAGTCA
16561	CAGTTTGACA	AACTCAACTA	CCCTGAGCCT	ATAGAGTGGT	AATAATTGCC	CTACTCATAA
16621	AGATGGGGTG	AAGATTAAAT	GAAATAGCAC	CTATAGAACA	CTAGTTCCAG	ACGTGGTATC
16681	ATGCTAGTAA	AATGGCTGCA	CAGCACTGCT	CAATGATGAC	AAAAAGTGAA	GCTTCTGGAG
16741	ACAGACTCCA	AGTTTGACTC	CCAGATCACC	ACATATAAGA	TGTGGGACTC	TGAGGCAGGT
16801	CATTTAATCT	CTCTGTGCAT	TAGTATCCTT	CTCTATACCT	TTACAGTGAT	GGTAATAGCA
16861	CCTACCTTCT	AGAAGTATGT	GAAGATTAAA	GATCCTTAAT	GCATATAAAC	CACTGTGTTT
16921	ACTGCTGTTT	GACAAATTTT	ATTTATAACC	ATCTTTACGC	TCCTAAAAGG	ACTTGAAGCA
16981	GCTTATGACT	GAAGACTTTG	GTAGGAGTTG	GCCTTCTATA	AATTATAAGA	ATTTCATAAA
17041	TTATTTGATA	TGAAAATGCC	AGTTGATCAT	AGTATGTTTA	CCGGGGTCCA	ACAGGTTGAG
17101	AAAAAATACA	CTTTTTTTCC	CTGAACATAT	GAAATTAGCT	CTCTAGGCAT	ATTCCTAAGG
17161	ACTTAAAGAA	TGATAACTAT	CATTTCTCTT	AAATCTTCCA	GATTTGGAAG	GATATATATA
17221	TTCAGCACAT	TGACAGACAA	TCCCAGTAGT	CCTAAATTAA	AAGACATTAA	AAATTAGTGA
17281	AACTTTTCCT	ACCTTTAGCC	TGTGTAATCC	TGGATGACCA	AGCATAAAAT	TAAATTGAGT
17341	AGAGTATACC	ACTGTAACAT	TTCCTGAAAG	GTATTCTAGG	CTCTGAGTAA	TTTCTTTGGG
17401	GTCTGAAGAT	CAGTTTGACA	TATCCTCAAG	TATCATGAGT	TCATTATAAT	TAAGAAAAAG
17461	AGAGTAAATC	TGGAGAATGA	GCCACTTTCT	TACTACTCCT	TGACCTCAGT	TOTOTOTO
17521	AGAGACAGGG	TCTCACTTTG	TTGCCCAGGC	TGCCAGGCTG	GAGTGTAGTG	GCGCAATCGC
17581	ATCTCATTGT	AACCTCCACC	TTCTGGGCTG	AAGCCATCCT	CCTGCCTCAG	CATCCTGAGT
17641	ATCTGGAACC	ACAGCAGGTG	CACACCACCA	TGCCAAGCTA	ATTTTTTAAA	AAGTTTTTTG
17701	TAGAGATGGG	GTCTTACTAT	GTTGCCCAGG	CTGGTCTCAA	ACTCCTGGGC	TTARCTCATC
17761	CTCCTGCCTC	AGCCTCCCAA	ATTGTTGGGA	TTACTAGTGT	GAGTCACTGT	ACCCCCCCCC
17821	ACTTCAGTTC	TGAGGAGGAA	AAAATATGTA	ATAATAATGG	GACTTTGGTT	TGCTGATTTA
17881	AAGATTCATG	TAACCTTATC	ATCCAATGCG	CAATTTGTAG	AATAATTAAT	AGAGACATOT
17941	GGTCTCATGT	TTCTACAGTT	GCTCATGCCT	TGATAGTAGA	TCTCCTTGCT	GCTGGCTCAG
18001	AAGGGTAAAA	GAGCAGAAAT	GATGGGGCTT	CTCTCATTCT	ATGAGGAAAT	AGACCTATGT
18061	AGAGGAGGCT	ACCTGTGGTA	AAACCTTATC	CTCATCACTT	AAAATTCTAG	GCTTATTCTC
18121	TGACCATATC	AAGTTTTCAA	ATGGTAAAAG	AATTGGATTC	AAGAGAAATA	TGAATAAACT
18181	TTTGTTTTCA	CTTTTCTCCC	TCCTCTCCCC	CCATTCTCCC	TTCCTTTATT	TTCTTGTCCT
18241	TAGTTTTCTT	TTCACTTTTT	TGTCTACTAT	TATTTGCCCA	AACTCAACTG	TAGGCTAGAA
18301	CAAAAAAAA	TTGAAAATTA	AAATGTGCCC	CTTTTGTTGT	TAGACTTGCT	TARACAATTG
18361	GGGTAATGAA	CCTTGGACAC	TAGATTTTAA	AACACACACA	TTTGAGCTTC	AGTGCACTGA
18421	AATAAATATA	TTTTTTAACAA	TTAAAAAATA	AAATTGCATG	TTTAAAAAAT	CTGCAGAGAA
18481	CAATACACGT	TGTGAGATCT	TGAATGGAAG	GAAAACTGCT	AGCCTCAAGA	GTGGATCAAA
18541	GATGCTCAGC	AGGCAACAGA	GTAAGAGCAT	GTTGGAGGGT	TTAGAGAGTG	TGCTCAGGGT
18601	TCTAGGCTCT	AAAAATCAGA	CAGTCCCCAC	GGCCTGGCCT	TCGTCGCTGT	ATCTTCTTTA
18661	TGAAAAACAC	TAAGTCTTTT	TCCTCACTGG	ATAAATTTTT	ATCCTTCAAG	TTTAGATCAA
18721	ATGGAACTTT	AGGACACTGA	CTAGGTTACA	TTCATCTTTT	AAGAGCGTAC	AGACATTCAA
18781	GGGCTAGAGG	ATGTGGGTTT	ACTGCACAGG	CTCATTATCC	AACAGCTGTG	CTACCTGGGA
18841	AACTTAACCT	CTCTGTGCCT	TAATTTCCTC	ATCTATAACG	CAGGGAGAAT	GACAGTAGGT
18901	ATCTCATAAG	GTTGTTGGAA	CAACTAAATG	CATTGGTATC	TATTGTGTAA	AGTGCTTAAA
18961	ACACTGCCTG	GCACAGAGCA	AACATCCAGT	GAACTTTAGC	CATCATCATT	ATCATTGTTC
19021	TCAGAGTCAA	ATACAATATC	TCATATCTGA	TAAATTACAG	AAGTGAATCA	ATCACTCTCT
19081	CTCTTTTCTC	CAGGGGGAGA	CAACAGCTTT	TAGACATATC	TTTTCCAACA	GTCGTCACTG
19141	CTGGACACTG	TTTCATCTTG	CAAATAAACC	AATGAAAATG	AGTGATCCTA	GAAGAAGATA
19201	AATGGAGGTA	TTTTGAACAA	TCAAAGAAGG	ACAAATGAAC	ACCTGGCTGA	GAAAAATTAG
19261	CTCTTTTTTC	TATGCATAAA	ACTATTAAAA	TATTCTTCAT	AGAAATTTAT	GACACAGGAA
19321	ACATAAAGAC	AAAATTAAAA	TAACTCCTAG	TATCTCCTAT	TCTTTTTATA	TGTATATTAT
19381	ATATACTCAT	ATTCATATAT	ACATATATCT	CACATCATGT	ATCATATATA	TTTAAATTAAA

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19441	AGGTGTCATG	ATATATATT	AGATAAATAT	ACTTAGAAAC	TTTTTTATGG	ATGTATAATT
19501	TATGGATATA	TTGATAATTA	TGTATTTGTT	ATTGACTACT	TCAATTGATT	CCCATTTTTA
19561	TGCATTATAT	TATAGATTAT	ATAGCTCACA	CATCTTTGTA	CATAAATCTT	TGTTCAAATA
19621	TTATTTCCTA	AGGATAGACT	TCATGAAGTG	GAAATACTAA	ATCAAAAGTG	AAAAACATTT
19681				TTGCTATTCA		
19741				AGCACTCATA		
19801	TCACTGTTAA	CCTAATAGTC	CTTCAAAAGA	AAAAAAATT	GAAATTACAT	TATTTTAATG
19861	ACTCTATTAG	TGAGGGTCAT	TCTTCCCATG	TTTCTTGTTA	GCCATGACCC	TATALGAAAT
19921	AAACTGCACT	GCAAAATGAT	AAACATGACA	TCAATCATTA	CATGGGAAGG	CACTATATAA
19981						CTGCGGAGGA
20041	CTCTGAAGGG	ATACTAAACT	GCATTTAGCT	GCATGCAACT	GAAACTACTT	TTACCTACAT
20101	TGTCTCTTAT	AAACATTATA	ACTACTCTTT	GAGAAAGTGT	TTACTATGGA	CTGAATTGTC
20161	TCCCCATCCC	CCCAAATTCA	TATATTGAAG	CCATAAACCC	CAATATGACT	CTATTCCTAG
20221				ATGAGGTCAT		
20281	ATAGGATTGG	TGGCCTTATA	AGAAGAGGAA	GATTCTGCAC	TTGGTCTTCC	מייממתדמממ
20341	ATTTATTTAA	AAGAAAAAA	AAAAAGAGGA	AGAGAGGGAG	CTCTGCACAT	ATACTCACCA
20401	AAGGCTATGT	GAGCTCTCAC	AGTGAGAAGG	TAGCACTCTA	CAAGCCAGCA	ACACACCCCT
20461	CAACAGAATC	CAGCCATGCT	ATACCCTGCT	CTGAGACTTC	CARGCCAGCA	AGAGAGCCC1
20521	AAATTTTCTT	GTTTAAACCA	CACAATCTAT	GGTATTTTTT	TATCCCAC	CARCCCARCA
20581	AAGACAGCAT	CATTGCTGTC	ACTUACACAC	AAGAAAACTA	ACACTACCAC	CAAGCCAACA
20641				ACAAGTGAGG		
20701	CCTCAATCCA	AGGCCAGGAC	TCCTCCACTC	CACATGTAGA	TRACERCE	ACCITCITCI
20761				AGACCAAGAT		
20821	CCTCATCTTG	DATABATATO	AGICAGCAII	TTACCCATGT	DARACAM	GAGACAAATG
20881	AAACAAAAAT	GCAAAGTATG	TACAAAAACTA	TGTTTACCAC	TTA A CTC A CA	ATCTCATGAG
20941	GCTTAATGAT	DICCTINING	TOTTCCACCC	GTTTGTATAT	CTCCTCACA	GTGATAAAAA
21001				GAGAGAAAA		
21061				ATATGATATA		
21121				TTGAGCTGTT		
21181	ATGGGACAAC	TGCACATGAC	ACTCARARA	CTCAGTCTCA	CCCATGCACC	AGGCACTGTA
21241				ATATCAATTA		
21301				AAGGGATTTT		
21361	GCTAGGCTGT	TTTGTTGGG	CCTCCCACCA	GCTGTCTAGG	COCCCOALCO	TGGCAAGAAG
21421	CTTCTATCAT	CCTGTGTTAA	CCATCTTCCA	TGTATCTTTC	A A COMMON MOO	ATGCAGGTCT
21481	CATGTCTAGG	GGTCATATCT	ATGTTCCATC	CAGGAAAAAA	AACCTCATGG	TCATCTGCAG
21541	AGGCATGTAC	CATTITANTO	CACACCTTCC	TTTTCAGAAA	GGGTAAAGGG	AAAGGGAAGT
21601				TTACAACGCA		
21661	TCTAATCTTT	TOACIATICE	CCTTTCAAAA	ACTGACTCAT	ACAGAAACGT	CACCTTAAAT
21721	AAAATTATTT	CAGTCATCCA	CTARTCACCT	GTTCATAGAA	TAACCTCCAC	GTGGCTTGGA
21781	GTGTTGTTAG	CATTATACAT	GIAAIGAGCI	GAATAAAAA	ATGTTTTGGA	CATCAAGTCT
21841	TCTTTACTTA	CATATAACTA	CTTANGCALL	TATAGCTGAA	CARCATGATG	TGGGTAAATT
21901	GTGGAACAGA	ADTABGATTA	CCTACATCOT	TCTCCTATGG	AAGAGAGGTT	GAAATGTCAG
21961	TCALACTE TO TE	CCCTCACCTA	CTCCCACAAG	TTCCTAATTA	GTGATTTTCA	GCTATGCTGA
22021	TTCCTCTCTC	CTCTTAGATA	CICCCAGAAC	TICCIAATIA	AATGGTGGCC	CTGATCTTAG
22081	AGCAGAAACC	TACTCAACAA	ATTATTO	CTACAGAAGA	TGTGCAGTTT	ATAAATGAGT
22141	ריים וויים ריים ריים	TCACTCACAA	ATTATTCAGG	CTCATCTGAA	CAGAGAGGAC	ACCTTCTCTG
22201	ACCACATAAT	AATTCTTCTC	CCCTGCCTTG	GGGTCAATTA	TIGICITGGA	CATTGATTTA
22261	AATTCTTAL	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	ALIGUTTATG	TTTGGATTTC	ATCTCCCAAA	ATAGATGGTA
22321		TOTOTOTOTO	CTCTARTACT	TAAAAAAAA	TITTGTGTGT	GTGTGTGTGT
22381	TLTTCIGIG	CARACCE	DAMPARORE	ATCGTACTTA	CACTTGTTAG	ATTTTTAGAG
22441	TACTCAGTAG	TTCAACCAAA	AATTATUTAU	ATACCCTTTC	TACAAAACAG	ACAAATTAAA
22501	TTTGAACACA	CTTABACCAAA	AAAAGCAGTT	CAAATAAAAT	ACTTGAAAAT	GAAGAAATCA
	ATADAGTTCA	ANATACTCO	ATCGTAAAAT	AATGTCTGTA	AAAATTATTG	CCAATCAAAT
22621	TABLEMENT	ACAMAMOACC	1 I GAAAAAGG	AAGAATCATA	TGAAAAGGGA	CTACTCATTT
		DOWINICHOR	MAMAGCCAAG	AAGTGAGTAT	GGTAAGAGTG	CTGTCAAGTG

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22681	AAACCCTGCT	AATCTCACTG	AACATGTAAA	AATCTGTAGA	TGCCTTTATT	TTATTCACTC
22741	ACACACATAT	GTAGAAAGAG	AAATATATGG	TAAACATTAA	AAAAACCAAA	TTAGAATGTA
22801	AAATTAATAC	TTTAAAAAAT	GGGCTGTATA	CTTTTCTTAT	CACCGGAGAT	AAGAATTTAT
22861	TATTTTTAAA	ATAAAGTTAT	TTTCTCTGTG	ACTGTTTCCA	TGACTTTGCT	ACTTAGAAGT
22921	TAGAGATGCC	AAAGTTTATC	TAAGAAAATG	TTTATGGAAA	TATTATTTCA	ATAATGAATG
22981	TTTAGAAGAC	TGAATTTCCT	GACTGGGCGC	AGTGGCTCAT	GCCTGTAATC	CCAGCACTTT
23041	GAGAGGCTGA	AGAAGGAGGA	TCGCTTGAGT	CCGGGAGTTC	AAGAGCATCC	TGGGCAACAC
23101	AGCGAGACCC	TGCAGCAAAG	TAAAAAGAAA	AAAGAATTGA	AAAAGGAAGA	CTGAATTTCC
23161	TTTGGGCAAG	TCATGTGACA	TTCCTGTGCC	TCAGTTTCTT	CATCTATAAA	GTTAATTCCT
23221	ACATTTTTGG	GGAAGGGAGA	GAAAAACTTA	GGATAGTGAC	TGGCACAGAA	GAAGCACTAT
23281	ATACTATATA	TATGTGGATA	TCATTTGTTT	TTATGGTACC	ATTTTAGCTA	TCTAATGCAA
23341	AATATGAATC	TTTTTTTTCT	GGGTCTTAAA	TTATGGAATG	TAAGAATTTT	CTAAATTCTC
23401	TAATTCTGTG	TTAGTTTTAA	AGCAATGGAG	TAACGTATCT	GTCAACTTGT	AAATATAAGG
23461	ATCAACCTGA	TCCACAATTT	GACCCCTAGC	CACTAATATT	TAATAGTACA	ACACTCAGAA
23521	ATTATCAAAG	GTCAGAGAAG	CCAAACAAAT	GTAAAAACAT	ACAGGTGCTC	AGAAAGATGC
23581	ACCTGTAATC	TCTCTAAGGA	GAAATATTTT	CCAAACTGAG	TCACACCCTC	CTTTAGTGAG
23641	TTGTGGAATC	AATCTCATGA	TTTCCAACCT	ACTCTTCTTT	TANANATORA	CTAGTCCACA
23701	GTAGAATATA	CTARAGTGCT	GGTGCTTAAG	ATACTATTCT	TANAMASIGAS	AAAAAAAAA
23761	ATTTTTTTT	TTTGAGACAG	GGTCTCGCTC	TTGCCCAGGC	TCNACTCCAC	TGGCACAATC
23821	ATGCTCACTG	CAGCCTTGAC	CTCCTGGGCC	CARCTCATTC	TCCCACCTCA	GCCTTTTGAG
23881	TAACTGGGAC	CACAGGTACG	TGCCACCACA	CCCCCCTANT	TOCCACCICA	TAGAGACAGG
23941	GTCTTGCTAT	GTGCTTAGGC	TEGECTETETE	AACTCCTGGG	CTCTACTO	TAGAGACAGG
24001	CAGCCTCCCA	AATTTATGGG	מת משלכים ומושל	TCACCCACCC	TROCTOCO	GTTCCCTGAA
24061	Դմահահահանանական 	TTTCACCTCT	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	CTCTCTCTCTCT	AMOGOMANA	CAGAGAGACA
24121	GAGAGAAGA	AACTTTTCTA	TOTOLOCATAL	CAATCAGAAG	ATGGGTATAA	CAGAGAGACA
24181	CTTTTCTTC	ארכוווכוא	CANATOTACA	CTCTCTCCTT	TITGAAGTCT	TATCTTTTGG
24241	CAAGGTCTTT	GCCATTCTTC	TCACACTATA	GCAACAGACT	CCCARCACTG	TCCCCTTAGG
24301	CCTTCTCAAA	AATGATTGTT	TATGCAATAA	ATTOTANACTO	CCCAACTICT	CAACAATACA
24361	ACABATTOTO	TGCTTAAAA	CTTCCAATAA	CTGCCGGGCG	AAGACAACTA	CAACAATACA
24421	CCCAGCACTT	TGGAGGCAGA	GCCGGGCNGN	TCACTTGAGG	TCCCCACTOR	CGCATGTATT
24481	TGGCCAACAT	GATGAAACCC	CATCTCTACT	AAAAATACAA	1GGGGAGTTC	GAGACTAGCC
24541	GTGGGCGCCT	ATAATCCCAG	CTARTCCCA	GGCTGAGGCA	CORCARMOCC	AGGCATGGTG
24601	GAGGTGGAGG	TTGCACTGAG	CCARCATCAC	ACCATTGCAC	TOOL COMMO	CTGAACCTGG
24661	CAAAACTCTG	TCTCABACCA	AACCAAAAACA	AAACTTCTAA	TCCAGCCTGG	GCAACAAGAG
24721	CAAGTATTTG	GGGATCTTCA	CANATGGGGG	TTATGGAGTT	TATCTACCAA	ATGTTTCACA
24781	GCTCTGGCCA	CACTADACTC	ATTCACCATC	CCAGAAAGGC	CTCLCCTTTGCT	GAGACCCTAT
24841	TCTTATCTCC	AGGCCTCTCA	Chargeate	TTCCAGTAGA	CTCAGCCTTT	GTGAGCAAGC
24901	ACATTATTCC	ADCADCCCTT	TCCCCACACC	TATGCAGCCA	AGCTCAGGGG	AGCACACTGG
24961	TAATTAAGCA	ATTCAGAGAT	CAGGGTCTCC	CCAGGCTGGA	AATCTGCCAG	CTCAGTTAAT
25021	AGCTCCTGGG	CTCTAACTCA	TCCTCTTCAC	TCTACCCAGA	GIGCAGTAGC	TGCGACCTCA
25081	GCCACCACAC	CCACCTAATT	TCCICIICAG	TCAGTAGGGA	AGCIGGGACI	GCAGGCATGT
25141	ACTCCTGGCC	TCCAGCCTTC	CCSSCTCCTC	TLAGIAGGGA	CCAGGCCAAC	CTAGTCTTGA
25201	CAACCCGCCC	ACTICTOCCTIC	CARAGIGCIG	TAATTACAGG	CATGAATCAC	TGCGCCCAGC
25261	GGTTCCTACC	TCATCTTTTA	TACTOR A TOTAL	CTGTAGTTTC	TAGTAGGTTC	TTGAGTCTAG
25321	GGTTCCTACC	GGGCNGGCCG	ATACACCCCA	AGGGGAGGGA	CTGTGTCTGT	TTATCTGGGG
25381	ATGTAGGGGT	ACCCCTCATC	ALAGAGGGGA	CTTCAATTAA	TGAAACCAGA	AGCAAAACTC
25441	AGTTGAGGAC ATCTTGATAT	TACCCCAMCC	MUMUTGGCCT	GATTATGGCC	AATCTTACAT	AATGTGTGAG
25501	ATCTTGATAT	CACABCCCAM	CONTCOMORGO	ATTAMAAGC	TACAGGGACT	TGGGAGCACC
25561	TTTAATTACA	CCTCTCCAT	GIICCIGIGG	ATTATGATTT	ATTAGATTGC	ACATGCCTAA
25621	ATAAAGACAT GACAGCTAAG	AGATCTCTCTCT	TACTTCCCC	1 TCTATAAGC	ATCTTCTGAC	TCCGCAATTA
25681	GACAGCTAAG TGGCGTGAAT	VOLUME TO TO TO TO	CTCTTCCCTC	ACATATAA	ATAATTTTAA	ATAAAAATCA
	TGGCGTGAAT	ATABETOTICE TO ATABETOTICE	TCCCARACCGAT	TIGAAGCTAT	CCATTIGGAA	GACCACTCTG
25801	AAGAGATGAA TTTTGTTCCT	CTCCCTC3 3 TO	TUCARAGAT	TACTTATTAA	TTTACAAGGA	AAAGGGGAAG
25861	TTTTGTTCCT	CICCGIGMMI	AAACACACAAA	ATCGAGGGCT	TTCTCGAATA	GTTTTGGCAT
<del>-</del>	CCAGGGTCAT	MIIMA	AAAADADAAAA	GICATGTCAA	ATATGAATTT	CCGCAGATTA

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25921	TTCAGCACT	TA GACCCTGGG	A GATTCTGTA	A AGAGGGGTT	TO TOTAL	C AACTTTTCCG
25981	GGTAAAAC	A ACACAAATA	C TCCTCCTCC	A AGGGGGGGT	C CCCCTCCCT	C AACTTTTCCG A GGTGATGCAC
26041	CAATCACAC	C GCGCCCTAC	C CTATATAAC	a acceptace	G GCGGTGCCT	A GGTGATGCAC T TTCATGCTTT
26101	TCGCTGGTT	A TTACATCTT	C CCAMALCACA	G TTCTTATCT	C GCCCGGGTG	F TTCATGCTTT  CCTGCAGCTT
26161	CTGCCAGT	C TGGTGTAGC	C GCTATGGAG	g ligitatet	C TGAAACCGT(	GCTGCAGCTT  GGGAGGAAGC
26221	CGGCTGGCT	T GATAAGTGC	A AGTCGCAAA	G TECCCAR	C CAAGAAGCG	A GGGAGGAAGC AAGTTGATCA
26281	CCGAGGCCC	T TTCAGTGTC	A CAGGAACGA	C TACCURACC	T CTCTGTGTC	AAGTTGATCA CTCAAGAAGG
26341	CATTGGCCG	C TGCTGGCTA	C GACGTAGAG	D TAGGIAIGT	C TITGGTTGC	CTCAAGAAGG CTGTCCCTCA
26401	AGAGCTTAG	T GAACAAGGG	A ATCCTCCTC	ADARTARCA	G CCGCATCAA	CTGTCCCTCA TCCGGTTCCT
26461	TTAAGCTTA	G TAAGAAGGT	G ATTCCTAAA	T CTACCAGGG	G TACTGGTGCT	TCCGGTTCCT AAGTCAGTTT
26521	CTGCCAAGA	C CAAGAAGCT	G GTTTTTNTCC	CIACCAGAA	G CAAGGCTAA	AAGTCAGTTT ACTGCTAAAA
26581	CCAATAAGA	G AGCCAAGAA	G CCGAGAGCC	GGGACTCCA	A GTCACCAAAG	ACTGCTAAAA AGCGGGAGAA
26641	AGGCTAAAG	G AGCCAAGGG	T DAGCARCAGE	CAACTCCTA	A AACTGTTAGG	AGCGGGAGAA AGGGCTTCGA
26701	AGTCAAAAT	T GACCCAACA'	T CATCAACAG	AGAAGAGCC	C AGTGAAGGCA	AGGGCTTCGA
26761	GAGCTTTCC	G GGAGGCCAA	T TTCCARGII	ATGTTAGAA	A GGCCACATCI	AGGGCTTCGA AAGAAGTAAA
26821	TATTTTAAG	A TGGCGTAAC	TIGGAMAGA	CCCAAAGGC	r CTTTTAAGAG	CCACCCACAT
26881	TTGTGATGC	A GCTGAGTTG	A PARCOLLINA	GITTCTGTG	A CAGTTATCTA	TAGGTTTAAG
26941	AGACCATCC	T GGGCAACAT	A CCCACACTOR	GATTGGAGA	A TTAATTCAGG	CCAGGCTTCA
27001	CCACCGACC	G GTAACCGGT	COTTOTO	CATCTATAC	AGGGGTCCTC	ATTTCCCCGG
27061	GGGTGAGCG	מבאתכינטטזני	CUTGTCCATC	GCACGTTATO	AATTGAGCCG	CACAGCTGAG
27121	TAGATTOTO	TARCETARCE	ACTGAGCTCC	ACCGCCTGT	AGGTTAGCTG	CAGCATTAGA
27181	GCTCCTTGT	TAMBULCAM	CIGIATICIC	AATGGCACAT	GCAAGGGATC	TAGGTTTCAG
27241	CATTGCTCC	T ACARICIAA	GCCTGATGAT	CTGAGGTTG	AGCAGGGGATC	GTCCGGAAAT
27301	TCTTGTGTC	AGCCCCIGC	CCCCCTGGTC	CGTGGTATA	TTGTCTTACA	CAAAACGGTC
27361	GTTGGCACG	TOCOUTTO	AGACTACTGG	TTTTACAAA	AAGTAAATTA	GTCAAGCATG
27421	TGCTACCTC	CTCCTIAGIC	CUTGCACCCA	GGCGTTTAAG	GATACAGTGA	GCTATGATGG
27481	AAAAAAGTTI	CICCAGCCIC	GGTGACAGCG	AGTCAGACGT	TGTCTCAAAA	CTTAAAAAAA
27541	GGAACTAAA	ACTOTOROGOUS	AGGGCTTCTT	GTCAGAGACT	GCCGTATATC	TAGAGGTCCA
27601	TACATGTAAC	ACCATCTANAC	CCAATCCTGA	AAAGCTCGAT	GGTGCACTAG	AGGAGGCTTT
27661	ATTTGGCATC	CONTRACTOR	TICIGGAAAT	GCCAGTGTCA	GGGAAGGGAA	GTGGAGAGCA
27721	ULDCIALOCATIO	TOTOCTOTO	TIGCIGATAC	TITTTTTT	TTTAACACAA	GTACTACATT
27781	AGGGATCANT	. IGIGGIGICA	TIGIAACTAT	TGTTTCTTAA	TATGCTATCC	ACTGACTTCA
27841	GTTGTTGTTG	TTCTTCTTT	CAAGGTGTCC	CAGAATATGG	ATTAGGGGAG	TTTTTTTGTT
27901	TTCCATTCTC	TCTCTCTTTT	TCATCTATTC	ATTATCCTGT	AGCTGAAATT	TAGAATTTTC
27961	TGGAAATCGT	CCTTCCTTA	AGAAATAACA	AATTTGTAGG	TTATAGTTGT	TGCAAGAATC
28021	TTACGGTCNN	CTCCTTAT	TTCCGAAGTA	CTATTAGGTA	TATCAACAAA	AACACACATA
28081	ATGAATTACT	TTTNACTANCE	AATTATTTTA	ATATTATTGG	TCTAATACAA	TTGTAACCCT
28141	ACCATACCCA	ACACMONA	TATTTATGAA	AAGAATCTGT	AAGTTTCATC	AGACTACCAG
28201	CCCAATTACC	TTCTCTAAAA	ATTITAAGAA	TCCAAACCTT	AATGGAAATG	TTGGAGGCTG
28261		TICIGAMITIC	CACCTTCCTG	AATCACAAAC	UK KAMPALADAM	TOTO I OMORO
28321		CGITICICIT	TAAACAGACA	TACTOTATATO	نته لا مانتخانجان بالمل	mmma
28381	ATTTCGAGAA	ACTTTCA LCAT	AAATAACCAA	TGCTAATGTT	AGTCTACTTT	GGACCATGGT
28441	אייייים מיייים א	TCCACACC	AAGTCCCCTG	CAAAACTATG	CATTGCATTA	TTTCACATAC
28501		TOCAGACGGI	LCAATAGTAT	كالمستمليات لاتالك		
28561		TURNAMITAL	GICCTATAAT	CAAACCTTCT	7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	
28621		MATCHARCEMO	AGACCTTTCC	THECTHECT		mman
28681		WATTWCINGG	ATTGTGCAAA	AATATCCCTC	TOTAL COMO.	<b></b>
28741		CCIMMIGCI	GIGCCCAGCA	ATCCACTCTC	3 CC3 C3 mmom	63 MG3 63 63 6
28801		WILLIAM TARGE	CTCTCCCAGC	AGCTGGCCGG		
28861		GGCTTCCTGC	ACCTTTCCTC	منحسات لاشابلت	CONTRACTOR	
28921		UCUITAVOQII	MAMAACAAAA	TCAATAACTT	DATECTATIONS A	000000000
28981		VOVCCVCTTT	TTTGTTTTGT		ATCCACCOCC	~~~~~
29041		GIGCHGIGGC	ACAATCTCGG	アサピ る	COTOMOGOOMO .	~~~~~~
29101		COCICAGCCI	CCCACGTAGC	TGGGATTNCN	COTCCCCCCC	
<del></del>		GIAITTTTAG	TAGACGGGGT	TTCACCATCT	TGGCCAGGCT (	GTCTTGAAC

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29161	GCCAGACCT	C GTGATCCAC	CACCTTGGC	TACCAAACTO	CTGGGAATAC	AGGCGTGAGC
29221	CACCGCGCCC	GGACTTAGA	CACTTTGTT	TGGCCAATAC	GACAACAGCC	TAGAACCCT
29281	CCGCAAATGA	A GAGCTTGTC	CTAAAGATGO	TTTATTTACE	TAGCTGTGTG	CCGCATGACC
29341	CAAAAGGTG	I TAACCTTTG	TCAACACGCC	CCTCCAGCCC	TTCGGTTAAG	TOCABACTAC
29401	CATTCTTAGA	ATGCTCTAA	\ ATACATAATI	TTTTTTTTT	Lilledadadadadadada	ب لا تاستسلسلسلس
29461	GAGTCTCTCT	CTGTCTCCC	GGCTGGAGGG	GAGTGGCGCG	ATCTCGGCTC	ACTGCAATCT
29521	CTGCTTCCGG	GCTAGCTGGG	CCTACAGGTG	CAGACCACCA	CGCCCGGCTA	AGTTTTGTAT
29581	TTTTTTTGGT	AGAGGGGGTT	TCACCATTTT	GGCCAGGCTG	CTCTCCCATT	CTTGATCTCA
29641	AGTGATACAC	TAGCTTTGGC	CTCCCAAAGT	GCTGGGATTA	CACTCCTCAC	CTTGATCTCA
29701	CAGCAAAATG	CTTTTTGTGG	AGCCAATCAC	TTTATTAGCG	CTTACCTCTC	TATGCCTACT
29761	TTATGCTTTG	AAATTTTGTC	ACAGTGTGGC	CGGTCATGGC	DAACACAAATT	CATTCTTATG
29821	CAGGATGTCA	CGGTTATTTC	TGTCATCCAA	ACTCATTCTC	GCDDCGCATT	TCAGCTCTTT
29881	AAACGACTTT	GTGAGCGGCC	CTGAAAAGGG	CCTTTGGGTT	. Արարարարարարարարարարարարարարարարարարարա	TGTTTTTGA
29941	AGTTCTCAGG	AGACCGCGTA	TTCTTAGATT	CAGCCGCCGA	ACCCATACAC	AGTGCGCCCC
30001	TGACGTTTTA	GGGCATATAC	TACATCCATG	GCTGTGACAG	TTTTCCCCTTT	GGCGTGCTCC
30061	GTATAGGTGA	CGGCGTCTCG	AATAACGTTC	TCTAAGAAA	CCTTARCCA	ACCTCGAGTC
30121	TCCTCATAGA	TAAGACCGGA	AATGCGCTTG	ACCCCACCCC	CCTTAAGCAC	ACGGCGAATA
30181	GCCGGTTTTG	TAATGCCCTG	GATGTTATCC	CGGAGCACCT	TACCARCCAA	CTTAGCACCA
30241	CCCTTCCCCA	AGCCTTTTCC	GCCTTTGCCG	CGACCAGACA	TCATTCCTAT	CGCAGTGGAA
30301	GGTATGAACT	GAAACAGTTC	CTTAAATACA	AACTTGGCGG	ACCTGATTGA	CGCAGTGGAA
30361	AGTTGGCGCG	GTTTTTTTT	TTTTTCAAAT	TTGGTCACCA	AGTGGGTGGA	AAACAACAIG
30421	CTGTTTCATT	ATGGTTCATT	GTTTTGATTG	GCCAGTGACA	GCTTGCTCTT	TCTCCCACTO
30481	GAAGGGTGTT	TGCAAGTTGA	ATGCGCTGTA	TTCCTGTCAG	CTTAATGACG	CTARCOAGIG
30541	CCCCATTCCA	CATTTCTTTT	TATTTCCACT	TGCTAACTAA	TAAATTACGG	A A WA COMMAN
30601	TGGGGAACAT	ACAAATAATG	TTTAAAGGAG	GTCAGATTTA	TAGGTCAAGG	AATAGTTTAT
30661	CCCAATCATT	TTAATATTT	TATTTAAACC	ACCCATTIA	ATGGCCTTCT	GATTTACCCT
30721	CAAGGTATAA	GTTTGGCTAT	GAAGTTTCAC	TCCTAARCAC	CCTATGTTTT	CTGTGCTGGA
30781	AAAGGTAGCC	AAATAATTGC	AAATTAAAAC	CTCATARCTC	CAAACTTCTT	GGGAAGGCAA
30841	TTCCCTATCT	CGATTCAAAT	ATTTGTTGAA	TCACTCATTT	TTCTGCAAAA	CCTCGTCACT
30901	GACAGGGAAT	ATAAACTTAA	GTCTGGATAA	TATCTTTTCC	CGGGACGCTC	GTCTGAGAGA
30961	GCTGTGCCTG	TTTGCTGTGC	CTGARATTCC	AAACACTCTT	CCCTTCCCTC	COMMUNICATION
31021	CCCCTTTCAA	CTTGCTACAG	CTTTAGAGAA	AAGAACATTC	GTTTTGTACA	COTTTTTAAT
31081	AATTGAAGTG	TAGGGCTAAT	ACTTGATTAA	GGTCATTACA	AAATCTACAG	GTTGGGGATT
31141	TGGGAGGTTT	TTGTGATAAG	ATTATTGGTG	TTDANTARCA	GCTAATCCCC	GGTCTTCCTC
31201	AATAGAATAG	CAGAATTGGG	TCTGAATGTG	CTTTCAACAA	AGGGACTTCT	TIGAAAAATA
31261	TTTTATTCTT	AGCTTCCTGC	GGGAGCTTTC	CAGAATGCCC	ATAAGATCCA	CAATTCAAAA
31321	AAAAACAAAA	ACAACCCCAC	CCACCACTCT	CTGGTTTAATA	ATAAGATCCA	CTTTTGTTTA
31381	ATTTAGAATG	GGGCTGTGGC	CTGTGAGAGA	CATTATATAC	TAACCTCAGA	TATTGGGAAT
31441	TGAAGAGAAG	AAATCCAGGA	ATGGAGAAAA	AAGACCCAGG	AAAGGCCAGA	AMGGMGMAGA
31501	TGTCATATTG	TTTGTATCAC	TTCTGAAATA	ATTGATTACA	TTCTTCTGCC	ATGCTCTACA
31561	TTCTTAGGTT	CTTCCACTCA	CTGTCCACAT	GCCACAACAC	AGACCTTATA	ACTACACTOR
31621	TAGCTAGGAA	GAAATGTCAA	ACATTACAGA	GAAAAAATGC	AGAGTCTGAG	ACIAGAGACT
31681	AAACTCTGAA	ATCTCAACAT	GCCTTTTAAT	TCATGADAAT	AAAAAATATA	CCACCATATO
31741	CAATATGACA	ATTCTCTGAA	AACATACATC	ATGTGAACTA	CCCTGGAACA	CARCTACACA
31801	AGTGCCATCT	TCATTTTAAC	CAGAGGTCTA	GGATGCCTTT	CCTTTATTTT	CCCTATTATA
31861	TCATTTATAA	AACCCCATTT	TTATTTTGAT	ATTTTATTA	CTTTCTATTT	CCTCCTCCTA
31921	ATATCTCCTT	TCTAAACTTT	TCTCAATGAC	AGTGACTCAA	AAACAATGAA	TCTCACAACA
31981	AATATTTAAA	GGATCTGTAC	ATGTAGATAT	ATATATTTAA	AATGGATTOT	TCCACACACA
32041	GAAGAATTCA	GGCATACTCA	ATCTTATGGT	TAGGGAGAGA	TTAGGGTTCAG	TCCCCTCTCC
32101	TGTATGGCTT	CTCGTTCGCT	TTCCATTTCA	CCTTCCTCTC	ACCCATCAGA	TCDDDCTAAIC
32161	ICATTGAACA	AGAGACCTAA	GCCCTTCAGA	TTAAAACTCT	GCAAACAAGT	ТСТССТТСАС
32221	AGGATACATG	AAGCATTCAA	ACAAATAAAT	CTATGATATT	AATCAGAGGT	<b>ፕ</b> ል አጥር ጥ አጥር አ
32281	TATTAATCAG	AGGTTAATGC	AGTGGCTCAC	GGCTGTAATC	CCAGCACTTC	AGGAGGCTGA
32341	GTTGGGAGAA	TCGCTTGAGC	TCAGGAGTTC	AAGACCATTT	TGGGCAACIIC	ACCA ACTOTT
						UCCUMO1611

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32401	CATCTCTACT	TAAAAAAAA	TAACCAGAGG	TGTTATGAAA	ATATAAATTG	TCCAGAACTA
32461	CCCTCCACAA	ACTAACTCTC	TCAGAATATT	CGATATGAGG	AATGAAATAT	GGTGTGTGTG
32521	TGTGTGTGTG	TGTGTGTATG	TGTGTGTGTG	TGTGTGTGTA	TGCACCTATA	TATGGCACCT
32581	ATATATTCAA	CAAACAATTC	TGATAATTGG	CCAGGGTTGA	GAATGACTAG	CAGCCCAGCA
32641	TACACTATCA	GTTTTAAGTA	TATAATTGCG	CTTTAGTAAA	ATGTAAAGAA	ATCCCAGAGT
32701	AGAAATACTT	TTAAGCTATA	TTACAGGTGA	GAAAATGCAT	AAGTATAGTC	TCACCCAAGI
32761	TAGACTATGG	GGGCTTTATA	ATGTCACAAC	AGTTGTTTCC	AGGCATTTGG	GGACATCACC
32821	ACTGGTCTTG	GGCAAGAAAC	TCCTCTAGCC	AATGGCTGAT	TTATCTCACT	CCCATCTAAC
32881	GCTTCACTGC	ATTTCTCTTT	TTCAGCAACC	TAACTTATTT	AAAAATATCC	ATTOTACE
32941	TCATTTTTT	CTGAATTAAA	CTGTCAGTAC	CATTGGCACA	CCTTTGGTTC	CCTACCATAC
33001	CTGTGTCTCT	GCTGTGTTTT	TTTTTTACCT	CCACTCCTTA	CTTTTCTAGA	AAAAAAA
33061	TGCTTTTTCT	TTTCAGTTTA	AATTATTTCA	CAAAAAGTTT	TCTTGACTTG	CACTTCCTAC
33121	GCTTGCTGTC	CTTGTGTGGG	CACGCTCCCA	TABACACTAT	TAATACACTT	CACIICCIAG
33181	AAAATAAAGA	TATCTGGACA	GAAAATTTCT	TANGLETICA	TAAGATTTTA	CGAIIIGIIA
33241	TGTTTATTTT	TTTCCTAGAC	TGGAGTACAG	TGGCACCATG	ATGGCTCATG	CENCOCONO
33301	CTTCCCCGGG	CTCAAGTGAT	CCTCCCACCT	CAGCCTCCCA	AGTAGCTGGG	A CER CROCER
33361	TGCACAACCA	CACCTGACTA	ATTTTGTTTA	TTTCTTTCCA	TTGTTTTTTG	ACTACAGGTG
33421	TCGCTCTTGT	TGCCCAGGCT	GGAGTGCAAT	GGCGGGATCT	CGGCTCACCG	AGAIGGAGII
33481	CT'CCCAGGTT	CAAGCAATTC	TCCTGCCTCA	GCCTCCCGAG	TAGCTGGGAT	CAACCICIAC
33541	CATCACCACG	CCCAGCTAAT	TTTGTATTT	TACTACACAC	GGGGTTTCTC	CATCOTTCACC
33601	CTGGTCTGGA	ACTCCTGACC	TCAGGTGATC	TECCCECCTC	GGCCTCCCAA	ACTICATIONS
33661	TTACAGGCGT	GAGCCACCAC	CONTRACTOR	TARTOTOTOTA	TATTTTGTAG	AGIGCIGGGA
33721	TCCCTGTGTT	GTCCAGGCTG	CTCTTC3 5 TT	CCTCCCCTTT	AGTGATCTGC	AGATGGGCTT
33781	CTCCCDAAAT	GCTAGGGTTA	CTCCCCTCAC	CCIGGGCIIA	TGGCTGGAAA	CCACCTTGTC
33841	AACATTATCC	TCTCTTAAAC	CIGGCGIGAG	CCACCAGGTC	AAACATGAGA	GATAATTTCT
33901	TTCATTTCA	7 7 7 TOTOTIANAC	CARTACTOR	AAAATTTTAC	ACAGAAAGCA	GTAATTAAAT
33961	TACATTTCT	TTATTCCCII	DAMIACITIC	CONNECTO	ACAAAGAAAA	CAAAGTATTT
34021						
34081	TGARGTTGGT	TATTTAAAACA	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	TATACATAGT	AAAACTCTTG CCGGGTGCAG	GGGGCAGTAG
34141	CTGTAATCCC	ACCACTTTCC	CACCTCCACC	TGCCATTCAG	CCGGGTGCAG	TGGCTCATGA
34201	CCACCCTCAC	CARACTITICS	AAAGGTCGAGG	CAGGCTGATC	ACGAGGTCAG	GAGTTCAAGA
34261	TGGTGGTGTG	TCCCTCTACT	CCCLCCTGIC	CLCCACTAAAA	ATACAAACAT GGGCAGGAGA	TAGCTGGGCG
34321	CCTCCCACCC	CCACATTCCA	CCCAGCTACT	CAGGAGGCTG	GGGCAGGAGA	ATCGCTTGAA
34381	ACCCCAACCC	CCCTCTCCA	GIGAGCCGAG	ATCGCACCAC	CGCACTCCAG	CCTGGGAGAC
34441	DAGGGGATANA	TCD ACCATAN	AAAAAAAAAC	AAAAAAGTGC	CGTCATAGTG	ACTCAGTTTT
34501	CAAACTTATA	CCAAGGAIAI	TIAACTCAAT	AGACTACAGT	TAGCTAACGT	GACTTGCACT
34561	DANAMACATATA	TTACA ATCOM	CACACONA	CCTGCCCCTG	AAGTATGAAT	TAAAGACTCC
34621	ANAMITCITI	TIAGAAICII	CAGAGTAAAA	GCTAGAATTT	GATTTTTTTA	AATAATAAAA
34681	TCCATCCCCA	AATTTCTAAATC	TGGTGTATAA	AATAACTTGG	TGGATGATGC	TTCAAGGCTA
34741	TTACAAATTT	CCCCCCCCC	COMOCOMONO	AGAGAATAAA	TGAATATGTC	AATTCAAAAG
34801					CCTTTCGGAC	
34861	CARTARATAR	TORGETTEGG	AGTTCAAGAC	CAACCTGGGC	AACATAGCCA	GAACCCGTTT
34921	GTGGGDGGDT	CACTTCACCT	TGAGCCAGGC	GTGGTGGTCC	CAGCTACTCA	GTAGGCTGAG
34981	CACACCAGGG	TTCCTCTCTC	CAGGAGGTCG	AGACTGCAGT	GAGCCGTGAT	CGCAGTACTG
35041	TTTGGCTGGG	CCCCCTACCT	ACTGAGACCC	TGTCTCAACA	ACAACAAAAC	AAGTTAGAAA
35101	CCATCATTC	ACCTCACCAC	CACGCCTGTA	ATCCCAGCAC	TTTGGGAGGC	CAAAAAGGGC
35161	ACTARACTIO	CARARARARA	TICGAGACCA	GCCTGGCCAA	CATGGTGAAA	CTCCATCTCT
35221	TECENTE	ACCCACCAAA	AMUCCGTGCA	TGGTGGCATG	CGCCTGTAGT	CTCAGCCACT
35281	ATCATCCCAC	TGCATTCCAC	ATTGCTTGAA	CCCAGGAGGC	AGAGGTTGCA	GTGAGCCGAG
35341	VICTIGCCWC	ATCANOCAS	CCTGGGTGAT	AGAGTGAGAC	TCCATCTCGA	GAAAAAAAA
35401	AAAATTCTGT	TROCKERS	CAAAATATCC	TTAAATTTTA	AAATACATCT	GAAAGATATT
35461	TCAAAATATT	TTTTCCTCTCT	ATTATAGGGA	TCAGGCAAAT	TCTGAGATTC	CTTTTTCCCT
35521	GCAGCAAACA	CTCACAGTGC	TGCTGTTCCT	AAAAACATGG	TAACTGTTGC	CACACCGTAT
35581	GTTTCCTTGG	CAUTTETET	AGGTTGTGTA	GITGITATTC	CAGAATAGCT	AGAATAAAAA
	TCCAGCACAT	CALLITUTTC	AGCAAGTTAA	CTAACCTCTC	TGTGCCTTGG	TTTCATAACA

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35641	GCAACATAAG	CATAACAGA	TAGCAGCAAT	AGCTCCTACC	י דברריים יים א	GATTCTTTGG
35701	AGGAATTAAA	TTAAGATTCA	GAACACAGCC	TAATATCTAG	TABGTAATAA	TAATTGGCTA
35761	AAAAAATTTT	CTTAAGATTA	TATATATTCA	TGGGGTACAN	CTACAATTTT	GCTACATTAA
35821	TATATTGCAT	TGTGGTGAAA	TCAGGGCCTT	CAATCCATCC	CGGDDDDDD	AAGTTTTTGA
35881	AAAGATTTCT	GCCATGGAAA	ACTTTTAATG	TACABATTCA	TCCATCCAAC	AAATAGAAAA
35941	TATATAAGTA	TCAACTCCAA	ATCCACCATA	ТСТАТСТСТТ	CTACACCAAG	AACAATTACT
36001	CAGAAATAGA	ATGCTTGAGA	TACCAGAATG	CATGCATATC	א א א א א א א א א א א א א א א א א א א	TGCATGCAGG
36061	ATGTCAACGC	ATCCTAGGCT	TTCAAATAAA	ATTGTCATAC	יייסועעועענעעע	AATATTGTAG
36121	TAACATTCTA	CATGTTAGAG	TGTAGAAGTT	AATCGCTGAT	CCDDDDDDCC	AAAAGAACAC
36181	ATTATACCCA	AAGCCTACAG	AGAGAATCAC	AATTACAAAT	ATCACCCTCC	ATGTGAAAAT
36241	CTTTAATTTG	AAAGTCAGAA	ATATTTAAAT	GATAGTCATT	GTTAAATCAG	ATTGTGGTTT
36301	GAAAAAAAGT	TAGTTTAAAA	CTGAGTTTAT	GAAAAATTTG	GGGATTTTAG	AGACAGTGTT
36361	TIGTTTTTAA	ATGTGTGTGA	GTTTGTGAAG	AATGTTTTAT	פאדונואט	CAGTATTATA
36421	AGATGACATT	ATTATAATAC	AACATAAGAA	TTTTGGCCTG	TACCTCTCTCAC	CAGTATTATA
36481	TCACCTGCTG	TACTTGACTC	AATGATTATC	AGAGTGGTTT	CTTTTTCCCCCAG	TGTTGTGTTC
36541	CCAGTTCAGG	CAGCTCAGCA	ATGGCCTGTG	AGAGIGGIII	TTCDAATAGG	TGGTAAGTAG
36601	TTTCTTGTTT	GTTTTCTCAA	ATTTTCAGG	COTTTTO	11CAMAIAGC	TCCAGTGCAC
36661	GCCCCTCCAC	CCATTCTTTA	TTCCTTTACC	TTCACCAAAA	CCCTCTCTCTC	TGCATCTCTG
36721	GTCACCGGAC	CACCGTGGTA	CATTTACCTA	TGGCCACCAC	CTCTCAGCGC	TCTCTTTACT
36781	ACCATGGTTT	GTGAATGGTT	TTGCCAGAGG	TGANTANGAN	TTTTAAAACCCT	AGGTCTTTGA
36841	TTTTTCAAAT	GTAGTTGACC	TTAAGAATTT	DTCDDTDDDC	CCACAAAATGC	TAAGCTTAAA
36901	AAACACCGAA	AGAAAATGAG	GACTTAAAAT	TTCTTTTTTTT	CCAGAAAAAI	GGCCACAGTT
36961	GCTGATGTTT	AGTAAATGTG	TTACTCAAAT	CTCTAITAAA	AAAATTAACA	GGCCACAGTT
37021	AAATCTCAGC	CCAGGTGAAA	TAAAACCAAT	DTDDDDDDDD	TCCTTT CCTT	GIGITICITG
37081	TGTAACATAT	TCCTTATGAG	GTAGAAGAGT	AIMAMACAAA	TGCTTACCTA	ATAAATTAAT
37141	TATAGTAAGA	TATTANGAGA	CARACACT	GTCNTNTCCT	TTATAGCAGT	TTTGCTGGTA
37201	AAATAACCAA	TGTCTTACAA	CTTAGACGAC	AATCTCCCTA	CACTORACIO	ACACGATTAA
37261	TTCGGCTACC	ACAGTTGAAT	GAAAATATTC	CCTDDCCCCIA	BACTGAAGAA	ACACGATTAA
37321	AAAATAAATG	TCTCCAAAAT	GACAAAGCGA	TTNACTATA	AAIGIAAAGA	AATTAGAAGC
37381	TCAATAAAAT	CATGCAGTAT	ACAATACAAT	ATACATTAT	TANACTATAT	GCATTTTTAA
37441	TGCAACAATA	ATACTAACAG	GTAATAGACA	ACTTCTTAAT	ACTOTOTOTO	GCATTTTTAA
37501	AATAACAGCT	TTAATTGTAT	TCATTTTATA	COTTTTCTAC	ANTERCOCON	ARCACRO
37561	ACTTTTTTCT	ACATAACTTT	TCTAACCACA	DADADAGAAA	ATTCCTTTANA	AATCACATTT
37621	AGATATCTTT	GCTAAAATTT	AATGCCTAAA	CDBCDDDCTT	CTCXCCTCTX	MOMAGAGATG
37681	TGAAGCACCT	GCCCTTCAAG	ACAGAATGCT	TGTACCACAT	TTATCCACCC	AACTCCATCC
37741	AGTAACATAA	AGTAAACACA	TGCCATCTGG	ATATATATATAT	TAIGCAGCC	AAGTGCATGT
37801	GGCAGGGTGG	CTCACACCTG	TAATCTCAGC	ACTTTGGGAG	GCCGACCCAC	CCCCATCACC
37861	AGGTCAGGAG	AGTTCGAGAC	CAGCCTGGCC	AACATGGTGA	AACCCTCTCT	CTRCTRRRRR
37921	TACAAAAATT	AGCCGGGCAT	GGTGGTGCAC	GCCTGTAATC	CCAGCTACTT	CCCACCCTCA
37981	GACAGGAGAA	TCGCTTGAAC	CTGGGAGGCA	GAGGTTACAG	TGAGCCGAGA	TCATCCCATT
38041	GCACTCCAGC	CTGGGCAATA	GAGTCTCAAA	ΑΛΑΔΑΔΑΑΑ	ACACTCTTTT	CARCOCCALI
38101	AACTGATTTC	CCAGAATCTA	GCAATTCCTG	AATGTCCTGG	TTACATTTTT	THE THE PARTY OF T
38161	GCACCGGAAC	CCCAGTGGCT	CCATGGAAGG	ACCTGGGCAT	CCTCTAAGCC	ACTTCCTCCC
38221	TTCCATTATA	CCATCTCAAA	ATGAGAGAGC	TTACTCCACT	TCATTGAGGG	ACTIGGIGGC
38281	AGAGTTCTGA	CTCCAGAGGC	ACTGGCCTAG	GGAGGACACC	GTGTGTGAAG	CCCACCACC
38341	CCACTAGCTG	TCCCCACCAA	TTACAGTCCT	TGCGTAGGGT	CCADAGAAAT	GANTECCAAA
38401	GAGAGCAACA	GAGGAGCAAG	GGAGTCACAT	TCCAGGACCT	TCCTTCAGGG	ACTITIONANC
38461	GAAACATGAC	AGCTGAGGAT	CAGTTGGTTG	TTTTCTGCTG	TTCCCCTTCA	TGTGATTCAN
38521	GCTCACTCAG	AAGAAACACA	ATGAGACAAG	AGAAGAGCCA	TCTCCTTCCT	ጥርጥርጥይጥጥጥ
38581	TTCTAGGCAT	CTAAACTACT	GAATGTAGTG	GTGTCTGAGA	TGTATCANAC	GGTCAGITIA
38641	ACTGAGTTTG	AAACCTGTTT	CTATCACTGA	CAAACTATGA	GATACTCTAT	VCALCUSATIO
38701	CTTTTTTTTT	TCATTTTTTT	ATTTTTATTT	TTATTTTTT	GAGATGGAGT	CACTICACTIT
38761	CACCTAGGCT	GGAGTGCAGT	GGCGCAAACT	CGGCTCACTG	CAAGCTCTGC	CICUCICIGI
38821	CATGCCATTC	TCCTGCCTCA	GCCTTCCGAG	TAGCTGGGAC	TACAGGCGTC	TGCCAGGII

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38881	CCCAGCTAAT	TTTTTGTATT	TTTATTAGAG	ATGGGGTTTC	ACCATGTTAG	CCAGGATGGT
38941	CTCGATCTCC	TGACCTCGTG	ATCCACCCGC	TTTGGCCTCC	CAAAGTGCTG	GGATTACAGG
39001	CGTGAGCCAC	CGTGCCCGGC	CTACTTCACT	TTCTTCATTT	AAAAAAGAAA	TGGGGATAAT
39061	AGTACCTATC	TCATAGAATT	ATTGTAAGAA	GTGCATGCAG	TAATGCATGT	AAGTAGGTGC
39121	TCAGAAGAGT	CGGACACGAA	GTAAGTGCTT	TTATCATCCT	TATCATAATT	TTCATTATCA
39181	GAACAAGGAG	AGACCAGGTA	GAAAATTATT	GTGATTCTTC	AGGTCTGGAA	TACTACACTA
39241	GCATCCCAAA	TGAAGGCACC	ATTAAACTTT	GCAAATCTGT	ATCACACCTT	CATCCCAAGIA
39301	AGAAAAAACA	CCTCTTCACA	ACCCCTTTCA	AGATATTTGC	CTCCTACCTC	CMIGCCAATT
39361	CCATCATACT	ACCCACAGAT	AGCCATGATG	CTTTTTCTGG	GACACCTCCC	TOTALAAAACAC
39421	GTGCAGTGTA	CAGCCTTCAT	AGCTGTGCAA	CTCACATCAC	NATCACATION	TCTTCCATTC
39481	AAGGCTTGGT	GACAGATGAG	TTACTCCCTA	ACACAGAGAG	ACCAMMONA	AAGAATCCCC
39541	AACGGGTCCA	GAAAATGCAT	ACATACATCT	GTAAAAATCT	AGGATICAAA	GGAAAAGTTG
39601	CGTCCCAGGG	TTCABACCTT	TTCTCACATC	TTAAAATGAA	GGIAAGGITA	TGACTAGCCA
39661	TAAGGAGTCC	TOTTOTANA	ATACCASAIG	AAATGACATA	COMCUTANCE	CCCCCAAATT
39721	CGGAGGAAAT	GDAGGDAGCC	TOTAGAMATO	GCTTGAGGTT	GGTGTATGTC	TCTGAGGTGA
39781	GAGAGGTCAC	AGCTAGGGAT	CACCGGGATG	CAGGAACTCA	CATGAGAGAC	AGTTCCAGGG
39841	TTTTTGAGGA	AATGAACAGA	CACCOGCAIG	ATCAAGGAGT	GAAACCTAAA	TGGGGAAATC
39901	TTAGGTTCAA	CTCTCTCCCTC	AAAGGCIAAA	AGCTCATAAA	TCGTCAGGCA	ATTTCTATGT
39961	CTAGTTTTCT	CTCCTTTCCCA	CACTCACTCA	GCAGGCTGCG	TGCACTCCCT	CTTTGAGTCT
40021	GACGTAGTCC	CCCATCCCTC	CAGIGAGICI	GCAGGCTGCG	TGTCACTCAC	GTTCAGCTAA
40081	ACCTAGGCAC	CATCUTCCCTC	CICCIGIGGA	GACAAGAGAC	CCAGGAAAGA	GGCATCACAA
40141	TAGACCTCCC	CATCITICCI	CTTCTCTCTT	CCTTATTTTC	CTCATTCACC	CATCTCAATT
40201	CTTTCTAACC	COTOTOTO	TTTCAAGAAC	CATTATCTCT	CATCTGGAAA	TGCTTATTGG
40261	GGDGATCCTC	CTCCTCCTCA	CCTCTCATCT	AACTTCTTAA	CAACACATTC	ACCATATAAG
40321	TATCCCAAAA	GACCCTTTCCA	CTTAGGATCC	TTCAATGACA	CCCCAGTGAT	CATAACCCAA
40381	CACCACACAC	CTTCACCCCC	CTCTGTATGA	GCTGGCTTCT	TTCTGATTCT	CTTTTCCCTA
40441	GGGTTD ACCT	ACACTA ATTA	TAGAAATGCA	TAATTGGTGA	GTGATAGCTA	CGCAAACTCA
40501	ATCCTTCCTC	ACAGIAATIA	TTTCTAATCT	CCCAGTATGC	CITATACTCT	CCTACTTGGC
40561	TTCTCCTTCA	CGICIGIGIA	GACCTCCCAT	CATCTTCAAC	CTCACCTAAT	GGAATCCAĢC
40621	TTCTCCTTCA	AGATCCAGAA	GGCTATCTTG	ATCCCCAGCT	GAATGTGATC	ATTCTTTCCT
40621	A CAMPERA COM	AAGCATTIGC	TTCCTGCCTG	CTTTAGGACC	TCATGGGGTC	TTCTTTAACT
40741	CACTTCTCAA	CCTCAAATT	TCATTCCCTA	CCAGATTTGG	GTTCTGAGAA	TAGCCACAGT
40801	TOTONNON	TOTTCOMMON	CCTGTACTAC	CTTAAACAGC	TCTTGCAAAA	TAGTAGGTGC
40861	CCCAGGGAGG	TOTTOTTOMAT	TAGAGACTTT	CATTCTGGGG	AGAACCATTA	TTTTCTGTCT
40921	CACATCCCCT	CTCCCCCCCCC	CCCAAAGAAT	ATAAATGAGA	AAAATGCTTC	CCATGGATGC
40981	CAGAICCCCI	ACTCACTTCA	TCCCACTGTG	CCCTGGGGCA	GAGGTACTAA	GAGACTTCCC
41041	ABCABGATGA	CCACAAAAAA	ACCOTGCCTC	TTCCTTAATA	TTATGAACAA	AATTCCAATG
41101	GGTGAGGGGT	CURLARARA	AGCAATTCCA	CTGATGACTC	CAATGACTAG	GGTGCCAGAC
41161	CCCCTAACAA	CIAAAACAGA	AAAAGCAAGT	TAAAGCCTTT	GATTGCCACC	CTCAGCCCAC
41221	CONCCCCON	MAGAGCAGATC	CTCATCTCAC	TGCCATAATT	ACCTCCTCAG	GCACTCCTCT
41281	TGAGGGGGGG	AMOGRACION	CAGCTCCTGG	CTCTCATCAG	TCACATACCC	CAGATCACAA
41341	CCCCTACACC	AICCAGGCCT	GGGTGCTCCA	CCTGGCACGT	ATATCTCTGC	TCTTCCCCAG
41401	TACCTTCCAA	CAAGGTTATC	CAGCCCTGGT	AGGTCCCATC	CCCATTGGGC	AATACGTCTT
41461	TCTCCCCCTA	CTCCTTGGCA	TCCATTGGCT	GCTTATCCTT	CAGCCACTTC	ATGGTGATGT
41521	CTCTCACCTT	GIAGITCAAG	GCCCGACACC	GTAGAGTGGT	CACTGAAGAG	GTCACATGAT
41581	TCTTTACCTT	CACCAAAGGA	GGCACTTGAC	AGGAAAGAGG	AAGGATGAGG	AGAGGGGATC
41641	TOTALACCCI	TGCCAGGAAG	ACTGGAACTT	TCACTTCCTT	CTATAGGTTG	GAGGAAGGAA
41701	ATACCCTTT	CAGAAAAAA	CAAGCTACAG	GAGAGACACC	ATTTTGTGTC	CTAAGATTGG
41761	ACTCIMACAC	AGTGTCACTT	GGAGAGCAGT	CAGATCAGCT	TGTTCTCCTC	ACATGTAAAT
41821	CARARATET	GITACCCATG	TICTITGTTC	TGATAGATAA	AATTGCCCTT	TATGTGCATT
41881	DENDECCES	GAATACAGAT	GGTCAGTTTC	ACCTGGGTCA	ACCTAGGAGG	CATTGTTATA
41941	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	ANGGAMAGATA	GGTAGCTTCA	GTGATTATTG	CTATGTTCTA	TGAAAGAAAC
42001	ATTCATATCTA	AAGGATTCTT	CTACTCTGAT	AAGTGGCCTC	ACTTGATATT	TTGTCCTGGT
42061	CONCERNO	TAGCTGAGAT	CICTGAATTC	TCTTTTTTTT	TTTTTTTTT	TTTTTAAGAT
4200T	GGAGTCTCAC	TCTGCTGCCT	AGGCTGGAGT	GCAGTGGCGC	GATCTTGGCT	CAGTGCAACT

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42121	TCCGCTTCCC	AGGTTCAAGC	GATGCTCCTG	CCTCAGCCTT	CCAATTAGCT	GGGACTACAG
42181	GTGCGCATGA	CTGTGACCAG	CTAATTTTTG	TATTTTTTTA	GAGACGGGTT	TCACCATGTT
42241	GGTCAGGCTG	GTCTCAAACT	CCTGACCTTG	TGACCACCCG	CCTCGGCCTC	CCAAAGTGCT
42301	GGGATTACAG	GGGTGAGCCA	CCGTGCCCGG	CCTTGACATT	TCTGAATTTT	TAACAGGTAT
42361	AAATATACAA	AAGATTATTG	GTTAAATAAA	AAGCAAGGGC	CATAGACACT	TCCCTTTGAG
42421	CCATATGCAT	GGAGAAAAGA	AATTAAACCC	ATGACTTGTG	GCTGTCTCAT	ACATCTCAAT
42481	TATAAGGTAG	AGACTCTAGG	ATTGAGAAAG	TCCCTTCCCA	GAATTTGGAG	AGGCACACAG
42541	CCTCAGCCAC	CTCTGAAACT	CCAACCAGGG	ATTCCGTGCC	CTGCAACCTC	CTCCACTCTC
42601	CCACTAGAGT	ATAGGGGCAG	AAGTGTGTTT	CCACCATACC	TTGTTGGTCC	AAAACACCTC
42661	TCCCCAGCTC	CAGCAACTGC	TGCAGCTGTG	CAGGGCAGTC	CCTCTCCAGG	TAGGCCCTGT
42721	TCTGCCTGGC	CCGAATCTTG	TGCCTTTCCC	ACTCCAGCTT	GGTGGGCCAG	GCCCTGGGTT
42781	CTGCTGCTCT	CCAATCCAGT	GTGTCAGGGC	AGAATTCAAG	GTGGTCCTGC	CCATCATACC
42841	CGTACTTCCA	GTAGCCCTCG	GTACTGTTGT	CTTCTTGCAT	TTCACAGCCC	AGGATGACCT
42901	GCAGGGTGTG	GGACTCTGGA	AAAATCCCCA	GCCTTGTTAA	CTGCAACCAA	AGGAATAGGT
42961	CCCTATTTCC	ACCATCCCCA	AGGACCAAAT	GATCTCAGGA	AGCAAATTCC	TTCCCTCTTC
43021	CCTGCTCCCA	CAAGACCTCA	GACTTCCAGC	TGTTTCCTTC	AAGATGCATG	AAAAGATGAA
43081	AAGCTCTGAC	AACCTCAGGA	AGGTGAGGCC	CCCTCTCCAC	ATACCCTTGC	TGTGGTTGTG
43141	ATTTTCCATA	ATAGTCCAGA	AGTCAACAGT	GAACATGTGA	TCCCACCCTT	TCAGACTCTG
43201	ACTCAGCTGC	AGCCACATCT	GGCTTGAAAT	TCTACTGGAA	ACCCATGGAG	TTCGGGGCTC
43261	CACACGGCGA	CTCTCATGAT	CATAGAACAC	GAACAGCTGG	TCATCCACGT	AGCCCAAAGC
43321	TTCAAACAAG	GAAAGACCAA	GGTCCTGCTC	TGAGGCACCC	ATGAAGAGGT	AGTGCAGAGA
43381	GTGTGAACCT	GGAGACAGAG	CAACAGGCCT	TAACCATGTG	TAGTAGGAGG	GGAGCAGGAT
43441	GTTGAGGCTC	CACACACCTG	CATCAACTCA	TACCATCAGC	TGTGTCTGGT	CCTCATTTTG
43501	TGAAGGGTGA	GTTGCAGTCC	TGTCTTTCTT	CCATATGACA	GTCCTGGGTG	CTCTTTCCTT
43561	GTGTGCTTTT	CTCTGCCACA	CGTGGCTGCC	ACCCCCTCAC	TGCCCCCAGA	TCCTATTCCA
43621	ATACTCATGA	TTAGACAGAC	TCCACTAAAG	CTGGTGGATT	CTAGAAAATG	TTAAGGTGTG
43681	TCTAGCCATG	GTAGTTGAAC	TCAGGAGTTG	GTGCTCAGGG	CAAATTAGAC	CCAAATCCTG
43741	AGGAATAATT	CCTTCAGTTT	TTTTTTTTT	TTTTTTTTT	TTTTTTGAGA	CAGAGTCTCA
43801	CTCTATCACC	CAGGCTGGAG	TGCAGTGGCA	CAATCTCAGC	TCACTGCAAC	CTGCACCTCC
43861	TGGGTTCAAG	GGATTCTCCT	ACCTAAGCCT	CCTGAAAACC	TGGGACTATA	GGCGTGCGCC
43921	ACCACACCAG	GCTAATTTTT	GTATTTTTAG	TAGACATGGG	GTTTCACCAT	GTTGGCCAAG
43981	CTTGTCTCAA	ACTCCTGACC	TCAAATGATC	TACCTGCCTC	AGCCACCAAA	GTGCTGGGAT
44041	TACAGAAGTG	AGCCACCGTG	CCCAGCCTTG	GTCCTGAATT	CTTACACTGA	ACTGCCTATG
44101	TGGCCTCACC	ACTTGGAAGC	CTGACTGGAA	TCTCAAACTT	AACATGTCCA	AATGCAGATC
44161	CTTGATTTAC	CCCAAACTGC	TCTTTCCTCT	GCCTTCACCA	TCTCAGAAAT	GGCATTGCCA
44221	ATTACCCCAC	TGCTCAGGCC	AATAAAATTA	AAATAAAGAA	CAAAGTCAAC	TTTAACTCTT
44281	CTCTTTTTCA	GGGGGTCAGG	GGAGACAGGG	TCTTGCTCTG	TCACCTAGGC	TGAAGTACAG
44341	TGGCACAGTC	ATGGCTCACT	GCAGCCTCAA	CTTCCTGGGC	TCAAGCAATA	CCCTCCACCT
44401	CAGCCTCCCG	AGTAGCTAGG	ATCACAGGTG	CATGCCACCA	CACCCAGCTA	ATTTTTGTAT
44461	TTTTTGTAGA	GAAGGGGTTT	TGCTGTGTTG	CCCAGGCTGG	TCTTGAACTC	CTGAGCTCAG
44521	GAATCTGCTC	TCCTTGGCCT	CCTCCTTGGC	ATGAGCTACT	ACACCCAGCC	AATTCTTCTC
44581	TTTCTCTCAC	ACAACATAGA	ATCCTTCAGC	AACTTCCTTC	AGAATATATT	CAGGAGACAA
44641	TGGTTTGTCA	CTCCCTTTTC	TGTTCCCACC	CAGCCCACTC	CACTACCTCT	TGCCTGGACT
44701	GTGTAACAGC	TTCCTGGCTG	GGCTCCCTGC	TTTTACTGTT	GCTCCCTTCA	TTCTGCTTTC
44761	CACATAGCAG	CCAGAGCAAT	CTTTTAAAAG	CCTGTGACAG	ATCACTGTTA	CTCCTTGGCT
44821	AGAATTCACA	CCACAGCCTA	CAGGCGCCTG	CACAACCTTG	TTTGTGGCTC	CTCTTCTGAG
44881	CCCATTACCT	ACTTCTTGGC	CTCTACTCCC	CAGCACTACT	TGTTTATTTT	TTTCAACCCG
44941	AGCTTCTTAA	CCAGGAGTTT	GTCTACTAGG	TGACATGTGG	CAAAGTTTAG	AGACATTTTT
45001	GGTTGTCAAG	ACTGGGGGAG	TGCTCCTAGC	ACCTAGTGAG	TAGGGAGGAC	AGGATACTGC
45061	TAGACATCCT	ACATGCAGAT	GGTAGTCCCC	CTTCCCACCC	CCACGCCGCC	cccccccc
45121	ACACACACAC	ACATGAGTAG	TGCTGAGAAA	ACCCGCTTTT	TAATCCAACT	TGCCAGGCCC
45181	ACTCAGTTTG	CCTGGGAAAT	ACTGCTCCCA	GTCAATATCA	TTCTTATTTC	CTTCATGTCT
45241	CTGCTCAAGT	GTCAGCCCCA	GAGTGACTTG	CCCTGACTTC	TCTGCTTCTC	ACAACACCCA
45301	TGATTTCCTG	ATGTTGTATA	TCTTTCTGCT	CATTTGCTTA	TTGTCATCTC	TCCCACTAGA

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45363	30003333					
45361	ATGCAAAATA	TCAAAGGGT	AAGACTTGTT	TCCCTGCTCT	CTCCCTTGGG	GCTTGAACAG
45421	TGCAACACAT	GGCTGGGACT	CATTTACACT	TGTAAACAAT	GAATATTTCT	GCTCAACATG
45481	AAATTTTATT	ATTCAACCT	TAATGCAGTG	TGATGTTTA	GAATCATAGO	TATGAAGTGG
45541	AGACATGAGO	: TCTGCCACC	AAGCCCAGTG	TACCATTGAA	TAAATTTGCC	AGGAAGCAGG
45601	CCGTGCCATG	CCTCATTCT	GTCATGTGTA	AAATGTGGAT	ACACGTAGTA	CCDDDDCTCD
45661	AAGTGCTGTG	CTGAGGCCGG	GTGTGACCC	ACAGAACACT	GTGCTACACT	ACAGGGCAAA
45721	ATCACTGTCA	ACTAAGATTA	L GAAGCAGCTG	TAGTACTTGA	AATAACATCA	GAAAACCAGA
45781	TTATTTATGT	TCTTTGTAAC	CTGAAAAGAG	TTATATAATC	TGAATTCCAG	מייים א כייייים א
45841	GTAAAATAAA	CGTATTATTA	GCTCCTACCT	CCCTATGCCT	AGTGAAAATC	AAATAAGATC
45901	AGATATGAAT	' GTAACTTAGA	<b>AGTGAGTGCA</b>	TTGCTTACAT	GTTCATTATC	YCLY CALLACA
45961	AGAGAGGCCT	' CTTAATTACA	CAGCACATTG	CAAATCAATA	AAGCCTAGCC	GAAAAGAGAA
46021	TTGTTCAGTT	' CAAACGTTCA	AAACTAACAT	ATACTTAATT	TTCCAGGCAA	AACAACAATT
46081	GCCAAGAGTG	GGGAAAGGCC	CGAGGTAGGC	CTCTCTCAGG	AGCCTCCCAC	CCTAGAGACC
46141	TCCACCCCAG	GTCTCACCAA	AAGTGGGTGG	AATGGTGAAG	AATTCAGATC	CCCAACGCCA
46201	CTCTTTCGCG	CCCCCACCGC	CCAACGCATT	CGTTCTGAGG	TGGAAACCCC	GTGCGGATCC
46261	TGCTGTGGGT	TTGCTCAGCC	TTCTCGGCAA	GCACTCAGGG	AAGAACTTCC	TGTTTGGAGA
46321	TGACTGGGGA	AAAAACTGCA	CAGCTGACAT	TGGAAATAAA	CCCGAGTTCC	AGGTTCAAGG
46381	AGCCCCAGGC	TTAGCTCAGC	TCAAGTGAGG	AACTACGAGA	TTTATTTAAA	AGCATTCTAG
46441	TTGGGGGAAG	GGAGTGGGCG	GTTCCAAAAG	TCACTCCGCA	GAGCCGGGAC	AGCCGGGGGA
46501	GGGGGCAGGT	CCTGGGGCGA	GGGACCCCTA	TCTGCAGTTC	AGTGGTAGGC	ACTCCCTCAC
46561	GGGGTCTGGA	CGCAGAAAGT	AGGGAGAGGG	GCTTGCGGAT	AGGGTTGAGC	AGGTCCTCCA
46621	AAGTTAGCAA	ACTCCCAAGC	GCAAAGAAA	AGCTAGTTTC	GATTTTTCCA	CCCCCGCCGC
46681	GCCCCTAGTT	CGCCCGCAGC	CCTCGGACTC	ACGCAGCAAG	CGCCCCTGCA	GGACCGCGGT
46741	CTGCAAAAGC	ATCAGGAGGA	GAAGCGCCGG	CCTGGCTCGC	GGGCCCATTT	CCCCACCCCC
46801	GGCCGCACGT	CCCCGTTAAA	TCTCCGCTTC	TTTTGGGGGG	CGGGGAAACG	CCCCAGCTCT
46861	CAGAAGTCAC	CCTACAGCTA	TTGCCTAGGC	TCAGGAGATG	CCCAGTAAAA	CTTCCTCCTC
46921	AAAAGCAACA	GGTCTTTCAG	AACTTTAGTT	CTCTCTCTCC	TACAGCAGAA	CCTACCTGGTG
46981	TGTGAAACAC	TAGGTGATCC	AGTGTCCCCC	TTGGTTTTA	AATCCTGAAG	CCCTCTTCTT
47041	GATTGGGGAA	AGTAGCTTCG	CAATGTTCTG	ATCTGAACTT	TAGATATTTA	AAMAMMAMA
47101	ATTTTCAAAA	TTCAATCATA	CATTTAAAAA	TTTTATCTCA	ACCTTAGACC	AAIAITIAIG
47161	TTATTTGACT	TAGAAATATA	AAGCTTTTTC	Valuation	TTGATTCAAA	TTT ATTT ATTT
47221	CATAACATTA	ACCAATTAGA	TCCTACTGAA	ACACCTTCCA	CAGCCTTCAT	1 I MAI I MAGI
47281	TCTGACAAGT	GTTTCACAAA	CTTTACAGTA	TTGGGATTAT	CTGGAGAATG	AATTGAATTA
47341	TTGAGGCCTG	CTCCTAACCC	CAGACACACT	CATTTAATCC	GTAATTGTTA	ATTAAACATA
47401	CATTAGCAGT	TGGGAGGGGA	TGACAGAAGA	GAGCGGAAAC	GCTGTCACTA	GGTAGTTAGA
47461	TGGCCCACCT	AAATTCAGGC	CCAAGACTAC	CCTAATCCCA	CCCTAAGGGA	AGACAGCCAC
47521	GATAAAGTCT	GTGGCCAAAA	TATCCTGGAG	AAACACAAAC	GAGGGTACAG	TGGAGTTTAT
47581	CCTAAGGTGG	CACATGCCCA	ACABCACADA	ACCOTOTOTO	CAAGTTCACC	GTGGAAATTC
47641	CATGCCATCA	TTATAATAGA	ATTTACATAC	AGTTTTGCCC	CCCCATCCCT	CCAAGTTCAT
47701	TCTTAACAAA	TTATAGGTAA	GACCATGCAC	ACTITATION	TAGATTGTAT	GGGAGGCTTT
47761	TTCAATCAAA	TAACATCATC	CTGTCACTCA	CATACACCCC	AAACCTCAAC	AGCTATACAC
47821	AAACCCCATA	AAAGCACCTT	GAGCTCTCTA	ANGANCECCE	GAGTTCACTT	TCCTCCCCAC
47881	AGCCCGCTGT	CCCTCAGAGT	GTATTATTGT	CCTTC A ATTA A	ACTTTGCTTT	CGCAGAAATA
47941	TTTGGTGTTA	GTTTGTAGTT	CTTTCCTCAC	TATCACARCA	ACTITIGCTTT	AAGCTTGCAT
48001	GCTCCGGCTA	TAATAATCTC	CTCGGTTAAA	CGATCCATGC	CAATGCATAA	CTGGTTCAGA
48061	CAGTATGGGA	TGCCACCTGG	GCAATGGGAT	TTTNNNNCCT	TTCCTTCTCC	TTCCCAGTAA
48121	TTTGGGAATT	ATTGCCTTAG	ACATTTCAAA	Chamamagui	AAATTTAATA	CTCAACGAAG
48181	GCTCCAAACC	TTTACATATC	TACCANATTC	ANCACCCAR	AAATTTAATA ATTTTTGTAA	CACCTGATTT
48241	AAATTTTGGC	AATTCAAGAA	DATCANALIC	CATATORCO	CCTCGACTGT	GCATGTATGC
48301	ATACAATAAC	ATTGGADACA	TCTACAARCAG	TCATCATCATCAT	CCTCGACTGT CACATTGGGG	AGGCAAACAG
48361	TATTCCTTTT	Thirty Value	TOIMGMAINT	TGATGATGGG	CACATTGGGG	CTGATAGTAC
48421	AAAATGCAAA	AATTGGCCCC	CCTCACTCC	TAATTAGCAT	ACCATATAAT	TCATCTATGT
48481	AAAATGCAAA CCGAGGAAGG	CAGATCACCT	GAGATCAGTGGC	CTTCCCCCTTGT	AATCCCAGCA	CTTTGGGCGG
48541	CCGAGGAAGG ACCCCGTCTT	TACTADADAT	DODA A A A REES	CCCCCCCCCCCC	AGCCTGGCCA	ACATGGTGAA
. –	ACCCCGTCTT		ACAMAMATIA	occessics I.G	ATAGCAGGCA	ACTGTAATCC

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# SUBSTITUTE SHEET (RULE 26)

48601	CAGCTACATT	AGAGGCTGAG	GCAGGAGAAT	CGCTTGAACC	CGGGAGGCGT	AGGTTGCAGT
48661	GAGCTAAGAT	CGTGCCATCA	CACTCCAGCA	TGGGAGACAA	GAGCAAGACT	TCATCTCAAA
48721	<b>Aaaaaaaa</b> a	TAGCTGGGTG	TGGTGGCATG	CACCTGTAAT	TCCAGCTACT	CGGGAAGCTG
48781	AGACAGGAGA	ATCGCTTGAA	CCTGGGAGGC	GGAGGTTGTG	GTGAGCCGAG	ATCATGCCAT
48841	TGCACTCCAG	CCTGGGCAAC	AAGAGCGAAA	CTCCGTCTCA	TAAAATAAAA	AAATAAAATA
48901	AAATGCAAAA	ATTAATGGAT	TTTAGTATAT	TTACAGAGAT	GTGCAACCAT	TACCAAAATT
48961	TTACATTTCI	ATCTCCCCAA	AAAGAAACCA	TGTTCCCCTA	ATTCAGTACC	CTTAATTCAT
49021	CGCCTCCCAG	ATTCCTCCAT	TCTCCTCCTC	CTCCCCTCCC	AGCCCTAGAC	AATCTTTAAT
49081	CTACTTTCTT	' TCTATTTGGA	ACATTTAGTA	TACATAGAGG	CATATAATAT	ATTGCTTTGC
49141	CGTGACTGGC	TTCTTTCATT	TAGCATAATG	TTTTTATGTA	TGTTTTTCAT	GGACCAATAA
49201	TATCTATTAT	' AAGGACATAC	CACAACATAT	TTTATTTATT	CATTCATCAG	CCGATGGACA
49261	TTGGTTTGTT	TCTACTTTAT	GGCTATTGGG	AATAGTGCTG	TTATAAACAT	TTATGTACAA
49321	GTTTTTTTGT	' AGACTTATGT	TTTGATTTCT	TTTGGTTATA	TATCTAGAAG	TGGGTTTGCT
49381	GGGTCATATG	GTAACACTGT	TTAACCTTTT	GAGGAATTGC	CACATTCTTT	TCCAAAGTAA
49441	GCATTTTATC	CTCCTATCAG	CAGTGTATGA	GAGTTCTGAT	TTCTCTCCAT	CTTTGCCTGG
49501	GTTTTTGAAT	CAGGGCCCCA	GATAGAACAA	AAATGTGGTT	ATTCAGTTGT	TCCACCATCA
49561	CTTGTTGAGA	AGACTCTTTT	TTCATTGAAG	TGTTTTGGCA	CCCTTATCAA	AAATCAATCT
49621	ACCATAAATG	TGAGAGTTTA	TTTCTGGAGT	CTCAATTTTA	TCCCATTATG	CTATAATCTA
49681	TAATCCTATC	TTTTTTTTT	TTTGACAGAG	CCTCACTCTA	TTGCCCAGGT	TGGAGTGCAG
49741	TGGCCCAATC	CCGGCCACTG	GCTCCTCCTC	CCAGGTTCAA	GCAATTCTCC	TGCCTCAGCC
49801	TCCCAAGCAG	CTGGGATTAC	AGGTACCTGC	CACCATGCCT	GGTTAATTTT	TGTATTTTTA
49861	GTAGAGACGG	GGTTTCACCA	TGTTGGTCAG	GCTGGTCTGG	AACTCCTGAC	CTCAGGTGAT
49921	CTGCCCACCT	CAGCCTCCCA	AAGTGCTGGG	ATTACAGGCA	TGAGCCACCA	CACCCAGACT
49981	ATAATCCTAT	CTTTATGTCA	GGACTACACT	GTCTTGATTA	CTATAGCTTT	TTAGTAAATT
50041	GAATTCAAGA	AGTTTCTCAA	CTTCAAATTT	GATCTTTTTT	TGGAAGACTA	TATTAGCTAT
50101	TCTCAGTCTG	CTGAATTTCC	CTAGGAATTT	TAGGATCTAT	TATCAATGTC	TATTCTATTT
50161	TTGTATATGT	TTTAATATTT	TCATAAGAAA	CTTTTTTCAT	TTAAACTTTT	TTTTTTAAGA
50221	AAAATAGTGA	AAATCAGAAC	ACTGGGGGTC	AGGCGCATTT	AACAGGCAGA	AGAAGAATAA
50281	AAACTTGTCA	TATAAACAAA	AAAGAAATGA	CCAATCACAT	TGTGGAAGCC	ATGGAGTGGT
50341	TATAGGTGCC	AAAGGCTGCA	GAGAAATGGT	GTCAGATATA	CCTGAAAATT	GTCCATTGTA
50401	TTTGGCCATT	AAGAGACTTA	GAAGACTTAA	GCCATAGATT	GCTCAGTGAG	ACCCCGAGGG
50461	CAAATGGTCT	GAAGGTGAAT	AGATCATTTC	<b>ACCTTTAAGA</b>	GAGCAGGTAG	GAAGCTATAA
50521	ATCCAAGATT	AAAAAGTTGA	CTGAACTGTT	AAGGAAGAAA	CTCTAATCTT	GAGCCACCCT
50581	ATCCTGGCTC	CACCTTCTGC	TGCAAGCAAA	CAGAAATGCT	GAAATTCAAC	ACTCACAAAG
50641	GCTGGTAAGC	TGGAAATGAC	AAAAATTACT	CCTGGGAAAG	TCAGATTTAG	AATTAGGCCA
50701	TATTTGTTGG	GGTTCAGATT	TTCATGTACA	CTTGGGAAAG	GGTTTAGCTT	ATAGGCACAT
50761	GCATGAAGGG	AACTGGTATA	GGGCTGTGTT	CATAAGGTCA	AGAGTTGAAG	GCCAGGCATG
50821	GAGGCTCTTG	CCTGTAATCC	CAGCACTTTG	GGAGGCCGAG	GCAGGAGGAT	GGCTTGAGCC
50881	CAGGAATTCA	AGACCAGCCT	GGGAAACATA	GGGAGATGCT	GTCTTCACAA	AACAATTAAA
50941 51001	AAATAAAATT	AGTCAGGTGT	GGTGGCACAC	ACTTGTGGTC	CCAGCCACTC	AGGAGGTTGG
51061	GAAGATCACT	TAAGCCTGGG	ACATTGAGGC	TGTAGTCAGC	CATGATAGTG	CTACTGCACA
	CCAGTCTAGG	TGACAGAATG	AGACCCTGTC	TCCAAAAAA	GAGCTGTATC	CACATCCCAG
51121 51181	GAAAGTGGTT	GAAGATCTAC	TTTTCTCTGT	AAACCTAATA	AAGAATAGAG	TGACAAATGT
51241	GTGTTGTGGA	AAGAAATGGG	GTGAGAGCTA	CGTAGATGCA	AAACAATACA	TCCCCACATA
51301	CCACTTGTTA	ATCATCCTTT	TCCACCCACT	TATGGGATGA	ATTGCATCTC	CCCAAAAGAT
51361	ACTUTGTOUT	AACCCTCAGT	AGCTGTGAAC	CTGACCTTAT	CTGGAATACG	GTGAGTTCAC
51421	1GGI IAAGAA	GAGATTATAG	TGGAATAGGG	TGAGTCCTCC	AACCAATGAC	TGGGGTCCTC
51421	DCD D D CCD P P	AGGGATGATG	GCCAGGTAGA	GATGGAGGCA	GAGATTGGAG	TTATGCTGCC
51541	ACCOUNTS	CACAGGAAGC	TGCTAGAAGT	GGAAACAGGC	AAGAAAGAAT	CCTTCCCCAG
51601	GAGAGARARA	GGGATCTTGG	CCCTGATAAT	ACCTTGATCT	CAACTGGCCT	ACGTAACTGT
51661	CTDACCARCE	ATTTCTTTTG	TTCTAAGCCA	CCCAGTTGAT	AGTACTTTGT	TACGGCAGCC
51721	TCTCTACCC	TGATATACAT	TTCTTTTACT	GTCATAGAAG	TTTTGAATCT	TTTAAGTAGG
51721 51781	AAAGGGGGGG	TCCTCCCAGT	GTCAACACAT	GGAATTCCTC	TCCTTGTGCC	TTGAAAAGTG
21/01	AAAGGTGTTT	GAACTGGTAA	TGAAAGAAAT	CTCAGCATGA	GGCCAGATGC	TGTACCTCAC

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51841	A COTOTA BETO	manacan amma	GGG3 GG3 BG3			
						TCAGGAGTTC
51901						AATGTTATCC
51961	TAGCCGGGCA	TGGTGCCTGT	AGTCCCAGCT	ACTCAGGAGG	CTGAGGCAGG	AGAATTGCTT
52021	GAACCCGGGA	GGTGGAGGTT	GCAGTGAACT	GAGATCACGC	CACTGCACTC	TAGCCTTGGT
52081						GCATTATAGA
52141	ATAAAAATGT	TTCCCCTTCC	CCCCAAACTT	TAAAAAAGCA	GAAGTCTGCA	TCATAAAATG
52201	GTCTTTGCCA	ATGTTATTTT	TATTATAACA	AAGGAATCTT	GCAAGGCTAC	CAGATCTCAG
52261	CAATTGTCAC	TATGTTCTGT	AAAAATCACT	TCCTAAAATG	TCTGAATTGA	CTGCTTGTCT
52321	CATTTATTTG	TTTCTCGTGT	CATACTGCAA	TGGATATCTG	TCTTGTTAGT	TTTATAAATA
52381						TATGAGGTAA
52441						AAAACCCCTC
52501	AGACACTGAG	TTAAAGAAGG	AAGGGCTTTA	TTCAGCTGGG	AGCTTTGGCA	AGACTCACAT
52561				TTCCTGTCCC		
52621						TGACTGGCAG
52681				GGATTTTCAC		
52741				GTCGGGGGTC		
52801				TCATTTCTGC		
52861	TTTCTTTGGA	GGCAGAAATT	GGGCATAAGA	CAATATGAGG	GGTGGTCGCC	TCACTTATTC
52921	ACCCCCTTTG	AGAATCTCAC	TCATTAGTGG	GAGTTCTCAC	TTTTATTCTC	ACTACCTATG
52981				ATATAGTACA		
53041						CAAACAAGGA
53101	AGCAGTAAGC	AGGTTTCTAT	TAATATTATA	ACTCCTATTA	TAAGAGTTTT	AAATCTTCTT
53161				CCAGAAACAA		
53221				ACTATGTCTT		
53281				AATTTCCTAC		
53341				TAGATAGCAT		
53401				TTATTAGTAA		
53461				GGGGTCCCAT		
53521				ATTCTCTCAG		
53581				TTTTTTTTT		
53641				CCTGTCTTTA		
53701				GCCCACTGGT		
53761				GGGGCTGTCC		
53821				TTACTAAATA		
53881				CTATTATACA		
53941				CCCACTTTTG		
54001				GGGTGAGTCT		
54061				GCTTCCCATG		
54121				GACATTGAGA		
54181				TGGAATTTCT		
54241				GGGAGCATTT		
54301	CCATATTTAC	TCAAGGATCC	AGTCCAGCCC	CAACTATTTC	TARGETTACA	CCTCAAACCA
54361	TTTTCCAGTG	AGAATCAAGG	GGGTTGGTTA	TTACTAGTTC	TANGGITACA	CACTCACCAC
54421				GAAGGTGGAC		
54481	ACCAAGTTGC	CTABATGACA	CARGACCAGT	ATCTACATTT	AGGALICILL	TIMITITIA
54541	CATGACAAGC	GTACTTATTT	TOTOCOATAT	AGCCTCTTTC	CTARTORAGE	AGICIIAAII
54601	CTATTTCTAA	CTTATTACTA	TTAATCACAC	CACAGGCATC	DARTHMONAG	GRANCEACATC
54661	TGGGCATTCC	Juliahalahahahahahahahahahahahahahahahahah	CTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	ACACTTTACT	CONTROCTOR	ATCARCTIGIT
54721	ACCAGTOOTO	AGTCCTCAAT	CITITOGCIA	AAACTGTGGT	COTATEGRA	TONONTOCOC
54781	CATANCACAC	ATCAGGTTGG	TATITIONA	GGCTACCTAC	CTTCTTTTTTTT	ATACCATTA
54841	ACAAACAAGT	TATTTTTACA	CALLICITA	ACTTATAATA	BCCBTARAGA	AIAGCATIAT
54901	TAGCAACTTT	TTGTCCTACC	TCACTCACTT	GATGTATACA	CTGGGAACAC	AAIAAGACIG
54961	GAGGAAGGTT	AGTTGAAGTC	TONGIGNOIT	DACTOCANAM	TTTT ACCAR	ARCAGREE
55021	TGATGAGTTT	TOTAL TOTAL	CCCCCyaccy	TGGACCAGTC	ACCERGGGAAA	MCMCAGTCCCT
			COCCATOCA	TAGMCCMGTC	AGC 11 CCGGG	IGIGACIGGA

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55081				GCAGGCGTTG		
55141				TGGAAGTTGG		
55201				GGGCTTTCTG		
55261				ATGCAGGGAT		
55321				TGATGTATTT		
55381	GAGGGCCTTT	AAGTTTAGGT	TTTGTCCAAG	AGTTAGCTTA	TCTGCCTCTT	GTGCTAGCAG
55441	GGCTGTTGCT	GCCAAGGCTC	TTAAGCATGG	AGGCCAACCC	TTAGAAACTC	CATCTAGTTG
55501	TTTGGAGGCC	CAGCCTCGGC	CAGGGCCCCA	CAGTCTGGGT	CAAAACTCCA	ACCGCCATTT
55561	TTTCTCTTTC	TGACACATAG	AGTGTAAAGG	GTTTTGTCAG	GTCAGGTAGC	CCCAGGGCTG
55621				GAAAAACTCA		
55681				CTGTCACCCA		
55741				GATCTGCTAT		
55801	CATCTAAATA	GATGTGAGAG	TTGAAAGACA	CATAAGGGTC	TTCTCTTGCT	TTACGATGTC
55861				CTAGGGTGAA		
55921				GAGTGCCCAG		
55981				CTGACTGATT		
56041				GGAATGCTAA		
56101				GAAGCTGGAT		
56161				AGAGCTTGGC		
56221				CCCATGCCTG		
56281				CTGCCATGTG		
56341			_	CATTCCAACT		
56401				CTTTAACTTC		
56461				GGTTGGGGCA		
56521				TAAAAGCTCT		
56581				CTCTGGAATA		
56641				AGGTACTITT		<del>-</del>
56701				AGGCAAGCAT		
56761 ·				TGGTCAGAAA		
56821				TTTCTTAGCT		
56881				GGTGGCACTT		
56941				GTCTCGGTGG		
57001				TTCCACCCTT		
57061				GGTGGTACAT		
57121				AGTATATAAC		
57181				TTTATAGTCC		
57241				TTTAGACCAA		
57301				CCAGATAAGC		
57361				TAAAACTCTG		
57421				TAGACTTTTC		
57481				ACCCGTTGTG		
57541				TTTAGCCAAT		
57601				AACCTTCAAA		
57661				CCTCATAATC		
57721				GTTTATTCAA		
57781				TCTTACATAA		
57841				CGACATGCCT		
57901				TTATCTCAGG		
57961				TTATTTCCTT		
58021				TATGTCTGTG		
58081	CAAGATTAGA	AGTTACTATA	ATACATGTTA	CACTGTTAAC	TTTTACCADA	ChhhbyChhhh
58141				TTATCCTTTG		
58201	AGTCCAAATT	AACTTAGAAT	TGGTATAGAT	GGCTTTTTTT	Tultulululululululululululululululululul	TACCTGGGAG
58261				TTCCTCCTAG		
				CLAG		

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				•		
58321		ATTTAGATCC				
58381		ATCATCTTCT				
58441		TGAGGTTTCC				
58501		CAGATTGAAT				
58561		CCTTAATGCT				
58621		CACTGCGTTG				
58681		GGCAATTGCC				
58741		CCGTAGGGAT				
58801		TAATCACCTC				
58861		GCTTACAACT				
58921		ACGTGACTGG				
58981		ACAATGCTTT				
59041	AGGTCAAAGG	TCGATCTTTA	ACCAGACCCA	GGGTGCGGTG	CCGGGCTGTT	TGCCTGTGGA
59101		CCCTTTTAAT				
59161		GGGGTGGTCT				
59221		CAAATATTAA				
59281	GTAAAGATAT	TTCTGTGGGG	AAAACATTTG	TTCATTAGTT	ATCAGTTAAA	ATTCTGTGAA
59341	AAATAACCAC	TAGAGACCCT	AAAGTACCCA	GGGGCTAATA	ATAAGAAGGG	AGGAACACCC
59401	TCTCACTCCC	CACCGTTACC	TGCCCAGAAG	GGAAGAGGAA	GAGGGTGACT	CCAGGAGAGC
59461	TGTGGTCTCC	CCTCCCCATA	TGTCCACATA	TACCTGACCT	CCCCTCCCCA	AAATATATAC
59521	CCAATATCTC	TCCCATATAT	ACATATTTAT	CTGACCTCTC	CACATATGTA	TACCTAAACT
59581	TTCTCTATAT	ATCCACATAT	ACCTAACCCT	CTCACACACA	TATAGCTGAC	CTCCAGTGGA
59641		GAAGAGAGAA				
59701	TGAAAAACAA	AAACCACACA	CAGAAAAAA	AAACACACAC	AAAAAAGAAA	TTGATAAATT
59761	TGTTTGTGTC	AAAATTAAGA	ATTCCGGTTC	AATGAAGGAT	CCCATGGATA	AAGTTAAGAC
59821		GGATGGTAGA				
59881		GCCAAGAAGA				
59941		CATGCCAGTA				
60001		GGCTGGCCAA				
60061		GTGGTGGGTG				
60121		CAGGAGGCAG				
60181		TGTGAGACTC				
60241		ATATATATAT				
60301		TGGTAATGAA				
60361		ATTAGAGAGC				
60421		CACTGCTGAT				
60481		CCCATAAAGG				
60541		TAAAATGAAG				
60601		ACGTGGAAGA				
60661		TTGTTCCTGG				
60721		AGTAAATGGC				
60781	AGAGATTGAA	GATTGTTCCC	TGGTCTGGGA	CCCTGCAACT	GAATATGCAG	AAAAAAGTAC
60841	ACCCCGCCAC	CCCGCTTCCC	ATCTTTCCTA	CCTGATTAGA	ATAGCTTTTT	CAGAAAACGT
60901	TGGCCAGGGG	TTGTGGCTCA	CACCTGTAAT	CCCAGCACTT	TGGGAGGCTG	AGGCGGGCAG
60961	ATCATCTGAG	GTCAGAAGTT	CCAGACCAGC	CTGGCCAACA	TGGCGAAACC	CCATCTCTAC
61021	TAAAAATATA	AAAAATTAGC	AGGGCATGGT	GGCACACACC	TGTCATCCCA	GCTACTCGGG
61081	AGCCTGAGGC	AGGAGACTCA	CTTGAAGCAC	AGTGATGGAG	GTTGAAGTTA	GCTGAGATCT
61141	TGCCACTGCA	CTCCAGCCTG	GGCAACAGAG	TGACACTTTG	TCTCAACAAC	AACAACAAAA
61201	CCCACCAAAA	CTTTAAATCT	ACCTATGGCC	AAATGCCTGC	TAAAATGAGC	ACCCAAGAAG
61261	CAGTGTTCAG	GAAAGTCAGA	TGAATACCCT	AAAATTAGAT	GCAATGTTGG	CTGGTCACAG
61321	TGGCTCAGGC	CCTGTAATCC	CAATCCTTCT	TGGGAGGCCG	AGGCGACAGA	TCGCTTAAGC
61381	TCAGGAGATC	GAGACCAGTC	TGGACAACAT	GGTGAGACCG	TGTCTCTACA	AAAACGTACA
61441	AAAATGAGCT	GGGAGTGGTG	GCGCGCACCT	GTAGTCCCAG	CTACTCAGGA	AGCTGAGGTG
61501	GGAGGATCTC	TTGAACCCAG	AAGGCGGAGA	CTGCAGTGAG	CAGAGATCAT	GCCACTACAC

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61561	CCCAGCCTGG	ATGATAGAGC	CAGACCCCCA	TCTCCAGAAA	AAAAAAAAA	AGAGAGAGAG
61621	AGATGCAATA	TTTAGGGTTC	AACAAGACTG	AATTTCTGAC	TCCTTTCCCT	ACCTCTCCAG
61681	CATGTTAGAT	TCTGGGTCCT	TCATCCTAAC	CCCCTGTTCA	TGCCATAGCC	ACCCTGTGGT
61741	ACCAACTTTG	GAAGCCTGGA	TCTTCATCCC	CTCATGATAA	TGAGTGTCCC	ATCAGGTCTC
61801						TTCACCCAGC
61861						AGGCCTTCCT
61921						GGCAGCACCA
61981						GAAGAGCAGA
62041						CTCTCTCCCG
62101	AGATGTTAGC	TAACTCATGA	GAGCCAGAAA	CCAACTGCAG	GCTGGCCTCA	GGCACTTAGG
62161	TAGTGCTTCA	GCCTCAGCAG	TCCACATTCT	AGGAACCCTC	ATAATATGGG	TTGAAGTATG
62221	CATTCCCACA	AAAATAAAGT	TGTTGAAGTC	CTAACCACCA	GTACTGAAAT	GGGDADAGTT
62281	CCCTTGTCCC	GCTCGCATGG	CATGTGATAG	GAGTGTGGCT	AATTTCTTCA	GTGCCTGGCT
62341	GCTCAAACCT	CTAGGGGAAC	ATTANGACGG	GCAGGTTGTG	GGTCTCCAAC	CCCATGACCC
62401	CACCACAGTG	TCTAGGGTTG	AATGTTTACA	GCTCCTGAAG	CCACAGTGGG	TETETETTA
62461					TTAACCAACT	
62521					CCAGGTTCTT	
62581					TGCAAGGTTT	
62641					AGTGGGAAAG	
62701					CCAGTCAAAT	
62761					TGCTCCCCTG	
62821					TGCAACATTT	
62881					CTGGGGGTGG	
62941					AAGGGACCAT	
63001					TGGCCTTATT	
63061					CCCAACATAA	
63121					TAGAGAGAAA	
63181					CTGGGGATAC	
63241					AGGAATCAAA	
63301						
63361					TAAATATCTG	
63421					TAATACAGTA	
63481					CCTAACCCCC	
63541					CAAACTCATC	
63601					GGCCAGGTGC	
63661					TCACCTGAGG	
63721					AAAAATACAA	
63781					GCTGAGGCAG	
63841					GTACTCTAGC	
63901					GCCAGGTGCT	
63961					CGCTTTAGCC	
64021					AAATTTAAA	
64081	GIGIGGIGGI	ACACATCTGA	GGCTCCAGCT	ACTCTGGAGG	CTGAGGTGGG	AGGATTGCTT
64141	TCTCCCCABA	GGAGGCTGCA	GTGAGCCATT	GCTGTCCAGC	CTGGGCTACA	CGAGAACCTG
64201	AAGCCCTACA	AGGAGAAAAC	AGTGAGACCT	CTTTTTCTCT	CCTCCTTCTC	TCCACTGCCT
64261	AAGCCCTACA	AGCACAAAAA	GGACACCACA	TGAGCACATA	GTGAGAATGC	TGCTGCCACC
64321	COTTOTO	GAAGAGAGCG	TTCACCTAGA	AACTGAATTG	GCCAGCACCT	GGATCTTGGA
64321	CITCIGAGCT	COLOGRAPIE	TGAGAAAGTT	ATTTTTTTT	TAGCGACTAA	GTCTATAGTA
	ACTOGRACIO	GCAGCTCAAG	GTAACTAACA	TAGTAGAAGG	GATGAATTAT	GGAGATCACA
64441 64501	AGTCCACGCC	TCCAGAAAAA	GACTTCCCTA	AAAATTAGTC	TGAGCAAAAT	TCGAATGATG
64501 64561	AATTATTTT	AAGAACTTTT	AAGGGATCTG	ACAAGTTTGC	AAGAGCTAGA	GAATGCTTTA
64561	CAACGTGATA	ATAGAATGCT	CTGTGATGAC	AGAAATCTTT	CCACACTGTT	CAAAACTAGC
64621 64681	TACTGGCCAC	TIGTGACTAT	TGTGCACTTG	AAATGTGACT	GGTGTCTGAG	GAGCAGAATG
64681 64741	TTTAATTTTA	CTTAATTTTA	ATTCATTAÇA	ATAGCTACAT	GTAGCTAGGG	GCTACTGGAT
64741	TGAACAGCAC	AGCTCGAGTC	TTTTAGAGGG	AGACAGGACT	CACCAAGATG	GATGCTGGTG

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SUBSTITUTE SHEET (RULE 26)

64801						ACATCCTTCC
64861						ATGTTACCAA
64921						ATGGTACTTA
64981						CCTTAAACTT
65041						TTTTCCCATA
65101						AGGCTTTTGC
65161						GACATACTGG
65221						AATTCTTAAA
65281						AAGAATGCTA
65341						CTCAGCTATA
65401						AATACGTATA
65461						CTGGCCAGAC
65521						TTAGGTACAG
65581						ACTCCTGTTC
65641						TTTGCAGAGT
65701						TCACAGTCAA
65761						AACACAATAA
65821	CTCTAATTAT	AACTGATGAC	ACAAACTCAG	ATATCAGAAC	TCTAGAAATC	CCCTATAATT
65881	TTGGAACACA	CATTCACAGT	TTTCACTGAA	ATATGACCTG	AAGATCAAAT	ATCACCTTAT
65941	TTCAACAATC	CTATATAACT	AAACGTGTCA	AATGATCCTG	TTTACCTCTC	CTTTGGATAC
66001	TCCAGGGGCC	CTCTGTAGCA	TCCAAAAGTT	AGGGGTTAGC	AAAGACAATT	TTGAAGCTGT
66061	AAAGGCTCAA	AACACTTAAT	GAACCTCTAG	TCATATCTGT	TCTCTACTCA	CTAAATGCTA
66121	GTAGCACCTC	TCAGTTGTGG	CTAAGCTGGG	AGGATCTCTT	GAGCCTAGAA	GTTTGGGGAC
66181	GCAGTGAGCT	ATGATTATGC	CACTGCACTC	CAGCCTGGGC	AACAATGCAA	AATCCTGTCT
66241	CAAAAACAAA	AACAAAAAAC	AAATTGCCTA	TGCTGTGGTT	ATCTCACAAT	TAATAAAAAG
66301	GAAAAAAAAA	GTATGCAGTC	TTTGTAGGTC	CTTGGGGTTT	GTTGGAACTC	AGAAAACAAT
66361	ACCCCAAAAT	AAAGACCGCA	GAAGCCAAAG	TTTTTCTCTG	ATCTTCTCCT	GCCCTCCTGT
66421	CTCTGAGTCC	CATTCTCCCC	GGAGTCTAGC	CATAGAAATG	AGAATTCCTC	TTCCTCAAGT
66481	TAGGTCATAG	AAATCAAAAC	ACCTTTTCCC	CAGAGCCCAG	CCATAAAACC	TAAAAATATT
66541	ACTCTAACTT	TCCCTCTGTT	TTTCTGTGTA	AAAACTGGCC	ATAAAGAAAT	TATCTGAACT
66601	ACCTTATTTG					
66661	CTGCACAGAG	AGGCGAAGAA	GAATCTAGAC	AGACAGGCCT	TGCTGGGTTT	CCCTACTCTG
66721						AAAATAAAA
66781	ATGGACAATT					
66841	TTTATTAATA					
66901	TCCCAGCACT					
66961	TGGTGAAACC					
67021	GTAGTCCCAG					
67081	TTGCAGTGAG					
67141	TAAAAATAAT					
67201	TCCTATTAAT	CTTCCTCTTG	TCGGTGGTTT	TCAGCGACTC	TTCAGAGGCC	AAAGAGTAAG
67261					ACTCTCATTT	
67321	TAAAGTGGCT					
67381	TCTCCTTTGA					
67441	GTTGTTGTTT					
67501	TCCGCTCACT					
67561	GCAGCTGGGA					
67621	ACGGGGTTTC					
67681 .	CTCAGCCTCC					
67741	TTTTTGGTTT					
67801	TATCATTTCT					
67861	TCTTGTCTAT					
67921	GCCAGTCTCT	GTACTTGGCT	GCTCCAGGGT	CCTACTTCAG	TTTCCCAGCT	TCTCAGTACT
67981	GTCACTGTCA	ATTGTGGGTA	ATAATTATTT	TTGTCCACCA	AAAGACTCTG	TATGTGAATG

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SUBSTITUTE SHEET (RULE 26)

_						
68041	AGTTTTGAAA	TCTGCTGAGT	AATACAGTGT	CAACCCAGTT	AATGATTTGC	CGGGCGGCTT
68101	GATCAGGGGC	TGTCCAACTA	CCGGCATTTT	GATTTGGAGC	GTCATCTAGT	GTCTGAAAGC
68161	ACAAACAACA	TCCTACATTG	TAAATGCCTT	TGGCTACAGA	GATTGAAACC	AAAGCAAACC
68221						AAAAGTTAAA
68281						GACAATTAGA
68341	TATTTTCAAT	TTAATGAAAT	TTTTTTTTAG	TTCACAGATT	AATACACAAT	GGGGGAGGGT
68401	TCTTATTCTG	TTGGACTTTT	ACATAACCTC	CACTTTAGTG	CAGTCTGCTT	TATGGGGTCT
68461	TGTTTGAGGT	GTGTGTGTGT	TTAAGGGAAT	GTGGTTTACA	ATCAAAATAT	TGGGTTGCTC
68521	TTAGGCACAT	TGTAAAGTCA	CACACCTGTA	TTCTTATTGA	TACATAATGA	TTAATAACAT
68581	TATTATTACA	GCCTGATCAC	CATCATTATT	GATATATCTA	AATAATGAAT	TTTATAATTT
68641	TGCTTCCTGT	CAGGCAAGAG	CCAATTTCAG	TGCTACCATG	TTTGTATAGC	AGTATTTATG
68701	TCTGTCATCC	TCAGTCATTT	TACTTCACTT	GTTCTTAGCC	AAACGGCCGA	GAAGCGATGG
68761	TCATTTTACT	TCAAAAATGA	AAAGAATTAA	TATTTTTACG	TTTCCCTTAA	AGACCCTATG
68821	TTTAACCTCC	ACTCCTGGGT	AAAATGGTCT	AGTCCCTCCT	TTTCATATCA	TCTCTGATAT
68881	CTTTTGCACA	GCCACTATTA	CCTACCGTTT	TCTAGATCCC	TATTCTTCAA	ACACCACCAT
68941	GAAGGTAGAG	CCTGTCTGAA	TTATTTTCTT	GTCCCCTGAA	CTCAGTACAT	TGTTAGGCTT
69001	CTTGAAGATG	TTGATCAGTT	GTTTGTGGAG	TGAATGAATC	AGCTAGCATG	ATTTTTTTAG
69061	ACCACTGAGA	CAAGTGTCTA	AGACACTTGT	TCCTTCCCAT	GTTCTTGCCT	GCCTGTGCAA
69121	TCCATGCAGT	CTCATGGCTT	CCCAGTGCCT	CAGAATTATC	CCCTGTCAAA	CAGGCATTAT
69181	AATTTCTGTC	CACTGAAAAG	GACAAAAAAC	TAAGTGTATA	GCTAGAAGTT	AAAAATTACC
69241	GGCCAGGTAC	TGTGGCTCAC	TCCTGTTATT	CCAACATTTT	GGGAGGCTGA	GGCGGGCAGA
69301					GGCGACCCCG	
69361					GCCCCAGCTA	
69421					CAGAAAAATA	
69481	CTCTTTCAAG	AGTTCGTGGT	TTTGACTGCC	ACCTAGCGTA	CATCAGAAAA	ACCECATEAC
69541	ATAGGAAATG	CCTGTGACAG	AGGGGTAAGG	TGAGAGAGGT	TGATGAAGAA	TGTATTGAAG
69601	GAGTGAAAAC	GCTTCCATCC	CTCTACTTAC	TAAATATATT	AGTTAAGTAG	TTGGGGGCATA
69661					TATTCTGTTT	
69721					TCAGTCTGCA	
69781	GAGAAGTCTG	AATTCTCATT	CTCCATTTCC	TTATTGGCAA	CGTGAGAATG	ATTACAATCC
69841					TCCATATAAT	
69901	GAGGAAGGGT	TCAGTTAACT	GTCTGTATTA	ATATTACTGA	TAACAGTCAT	GDCDDDCDDD
69961	AGCTTAACAA	CAACACCACC	AACAACAGTT	GCAGAATTGA	GCCACCAATT	TGCACACAAA
70021	ATTGTAGGTA	GGATGTTTTA	GAAAAGTTAT	TATTTAATAT	ATGTATATAT	TTTTCTACTO
70081	AAAATATGTC	AGAGGTTGTT	CTAAGAACTA	TTTAAATGTT	AACTCCTTAA	TCCTCATAAT
70141	GACCCATGAA	ACAGGTAGGC	TTATTATTGT	СТСТТАСАТ	GTGAGAACAC	TGAGACACGA
70201	AAAGGTTTAT	TAACTCACCC	AAAGTCACAC	AGCTGGTAAA	ACGGCAAAAT	TGABACACGA
70261	CTCAGACATT	CCAGGTTCCA	AGACAGTCTA	ATTATTCTTT	TGACTAATAT	ACTARGCTGC
70321	CTCTGTATTT	TTCCTTGATT	ACTTTGTAAA	AGTATGAGGA	AAATATAAGT	GCTTCAAGTA
70381	ACCATGAAAA	ATATAAACAA	TCTATGTATC	AACTGAAGCA	TAATTACAAA	TCCTTTCATA
70441	AGCAAACATA	ATAAAAATTT	GATATCAATC	AAAACTTTCA	TGTAATGTAA	CCTTTGATA
70501	ATGAATTCTA	TAGTAAAAA	GTGCAGAGTG	CTGGAATACC	ATGCTCCTAA	TATATTCCCT
70561	AGGCACACCT	GCCTGCTATC	AAAGGTATGC	ACACACCTTG	GATACAGAAA	CTTCCCCACTC
70621	GGTAGTTATG	TGAGTGTCAT	CAGAATTCTT	TCCCACTTGG	GAAAGAATTG	TCCATCATA
70681	GCTTGGATGA	TGGACAAGGA	GTGAGCTCCC	AGAACAGTGA	TGTGGGGATA	CATCCTCACA
70741	TCACAGTGAG	AATGAGTGTT	CTAGACTGTT	TACACACCTA	CCACTCCTAA	ATGCACACAT
70801	ATAATTGCTT	GCACACACAC	ACATACACAC	TCATCTCTTC	TCTGGTGGTC	CAGCTCTATC
70861	TCTTATCATT	AGGCTTCTTG	GGGCTAGTAC	CTAGGGCCTG	TATCCTTTCA	GAGGCAGCTA
70921	AGGGAAGCAC	ACATAATTAG	AAAGAATGAA	CCAGCTTGTT	GGATTTGGTC	TOTOGONGCIA
70981	CAGCCCTCCA	AGTTAAGGAG	AGTACCATCT	TTCTTAGGGT	CACCAAAGGA	Λαπασαπιο
71041	AAAGAAAGAA	ACAGAAGGAT	ATCATACAGO	AAGGATCTAA	TGCDDDTDTG	CCTCDDDTCA
71101	GAGGCTACTG	TGTGCTGATC	CCAATCCCAG	GAACTGTATG	CPCPATATATA	ADTITATION
71161	TCACTGTATT	TCTGGGAGTA	TTATTCCCAT	TTTACACACA	PUCKTINICI	CAGGGTANCC
71221	AAGCTCATGA	ATGGAGAAAC	TGGGATTAAA	TATAAAACCTT	CCALCCACCY	CAUGGIANCE
					CCTIGCICCA	OWNCIRCIB!

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71281	CTTTCTGCTC	TTCCACACTA	CCAGCTCAGC	TGTGCTCTCT	ACATGCAGGC	AGTTTTACAA
71341	GTTTCAGATT	AGCCTGGGAC	TTCCAGGGTT	TTGAATGGGT	TAGGGAATGG	GGAACTTTTG
71401	GGTTTACTTT	CCATTTTTTC	TTCATACATA	TGTAATATAT	AACATAAATC	TATGGTATAT
71461	ATGATAAATA	TATGGCTACA	TATGAACTAT	ATAATCACAT	ATATGCATTA	TAAATAAATA
71521	TTAATTTTAT	AATATTTTAA	AGGTTATCAA	ATAAATATTA	ATATAAATAA	TTAAATAATT
71581		TTTGTTTTCC				
71641		TTTTTAGGAG				
71701		TTGTCTTCTG				
71761	ACGTTTAGCC	AGCACATTAA	CATTTTATGT	TTTTATTCTT	TTGTGCTCTC	AGTGGCTGTG
71821	TCTTTTCTAT	CGATTTCTCA	CACTGTATGA	TGGTTATATT	TGTCTGTATC	TGTCCCACCA
71881		CTTGAGAGGA				
71941	GTGTCCTGTG	CTTAACAAGT	GCTCATTAAG	TGTGTAAAAA	CACAGCACAG	TAAAAAACTA
72001		AATAATGTCA				
72061		TGTCAGGTTA				
72121		CTTTATAACT				
72181		CAATTGACAA				
72241	TTTGACATCA	TTTGAAAGAA	GCTTGAAGAA	TAAGATAGCT	GTTAATGACC	CAGTTTCCTA
72301		ACAATTATAA				
72361		TTGTAATACT				
72421	TATTTATTTT	TCTGGACAGA	GTCTTGCTCT	GTTGCCCAGG	TTAGAGTGAA	GTGTTGTGAT
72481		TGCAACTTCA				
72541		CGGGAACTAC				
72601		TGTGCTTTAG				
72661		ACAGATGCAG				
72721		ATCTAATTGT				
72781		TTATATGTTA				
72841	GCCCCTACAA	TCTTATTTAG	GCTGAAAATA	TCCTGGAGTC	CCTGTATTGT	TGGTCTCAAG
72901		CACTAACACT				
72961		ATTAATTGGT				
73021		ATTTAGTAAT				
73081		CTTGGTGGTT				
73141		GTGCCAAGAG				
73201		AAAGAACTCT				
73261	TGTCACCCTC	ATTCCTTACA	CCTGTCCTAA	CAACTCCTCT	CACTATCCTT	TGAATATATC
73321		GAGTCTCTCT				
73381		CTTTTCTACC				
73441		ATAATATTTA				
73501		TAACTGTGTC				
73561		CACATCCTTA				
73621	ATAGAGTCTC	GCTCTGTCAC	CCAGGCTGGA	GTGCAGTGGC	GCAATCTCAG	CTCACTGCAA
73681	CGTCCGCCTC	CTGGGTTCAA	GTGATTCTCC	TGCCTCAGCC	TTCCAAGTAG	CTGGGATTAC
73741		CACCATGCCG				
73801	TGTTGGTCAG	GCTGGTCTTA	AACTCCTGAC	CTCATGATCT	GCCCACCTCA	GCCTCCTAAA
73861	GTGCTGGGAT	TACAGGCGTG	AGCCACCCCG	CCCGGCCAGA	GGTCATTCTA	ATAGACTTTT
73921	TITTTGTTGT	TGCTCACAGG	CTTGTTCAAT	CTTATTTCAA	AATTTGAGAA	ATACAGTTTC
73981	CATGGAACAC	CAACCAGATA	TCAGGTTGCT	ATGGAGTTGA	TAGTCAAAAG	CTTTGTATCT
74041	TCCAGTTTTT	CAGAATGGCT	TCTAAAGGTT	CTGATTCAGA	GCTCTTAGGC	GAAATTGAAC
74101	AACCAAGTGT	CAAAGTACAA	CATTCAGGAA	GTTAAAAACA	TGACTGACAT	ATATGTACTA
74161	ACARAGEGA	GCTTGTGTAT	GTGTCAATGA	ATGATTTAAT	TCATTAATGA	AGGAGGAAGC
74221 74281	AGAATCACAA	TTAGGTCAAA	GGAAGATACG	GGAGAATAAA	ATATGTATTT	GGTCAGGGAA
74281	CONTROL	CTGGAAGAGG	AAGGGAAAAT	CAGATATAAA	GTTGTTTAAT	GACTTATTAG
74401	GCAATACAAT	AATAACTTIT	AGGGTCATTT	TITCTATATT	AAGAATTCAT	TTCCATCTCT
74461	ATGACAAAT	CCTTATTAAT	TTATTAAACT	TCTACAAGTG	AATGTTTACT	TTTAGATAGT
, 340T	CTGGACCCAA	TAAAATGTAA	ACATTAAGTC	AGAGTTACTT	TCACGTAGGA	CAGTGTTGTC

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74521	CAATAAGGTA	CCACTAGCT	A CACGTGATCA	TTGACCATT	GGACTATAGO	TAGACTGATT
74581	TAAAATGTTC	TAAAAGTGTA	AAATACACAC	CAGGTTCTG	AGATTTATCA	TTTAAAAAAAG
74641	AATGTCAACT	GTCTTTTTT	TTAGCTTATT	TATTATATG	TGAAGTGATA	ATAGTTTAGA
74701	TATATTAAGT	TAAATAAAT	ATCTTAAAAT	TAATTTTACT	TGTTTCTTT	CATTCTTTCA
74761	ATGTGACCAC	TAGAAATCT	GAAAGTATTT	ATGTGATTC	CATTCTATTT	TACTGTCTAG
74821	TATTGCCTTA	CATCATCAGG	TACCCCATAA	GTAGGCTTTT	TAGATAATTO	TCTAATATAG
74881	CTTGGAAGGA	TATGGAGAAA	TATTTTTGCG	TTGCTTTTA	GTTTTGCATA	ACTTTTTCAA
74941	CACACTTTAT	AAAGGATCTA	GAAAAGGGTT	GGTTACATGT	TTCTCTGTCT	TCTGGCCTCC
75001	ACCATGTTGC	CAGGAGGTTG	GGGACAAGAT	TCTGGGTGGC	TGGATGTCCT	' AATGGCTTGA
75061	GGTCTGGACT	TGAGATTTGC	ATATAAAGAG	ATGTGATTAG	ATTGAGTCGA	CTAGAAAAAT
75121	CATATTAGAG	AACTGAATCA	CAGCGATTAA	ATTTACATGT	CGATTTATAA	ACCAGGACAC
75181	CAATTTATAG	TGAAAGAAGG	TCCAGTTACC	TGGTAATCAA	GACGTTTCAT	AGCTATTTTC
75241	ATGATGGATA	TACTTAGCTG	AGTTTTAAAT	GAGAAGGGGG	TTCATTGCAC	ATAGAATAAG
75301	ATCTAAGTGA	. AATGTTTATT	TTATTTTTTT	TTTTTTGACA	TGGAGTCTTG	<u>רדרידנידיניר</u> ר
75361	CAGGCTGGAG	TGCAATGAGG	CAATCTCGGC	TTCTGGAGTG	CAATGAGGCA	ATCTCGGCTT
75421	CTGGAGTGCA	ACGAGGCAAT	CTCGGCTCAC	TGCAACCTCC	ACCTCCCGGG	ТТСАВАТСАТ
75481	TCTCCTGCCT	CAGTTTCCTG	AGTAGCTGGG	ATTAGAGTTG	CCTGCCACCA	CGCCAGGCTA
75541	ATTTTTGTAT	TTTTTTTAGT	AGAGATGGGG	TTTCACCATG	CTGGCCAGGC	TGGTCTCGAA
75601	CTCCTGACCT	CAGGCGATCT	GCCCGCCTCA	GCCTCCCAAA	GTGCTAGGAT	TACAGGCGTG
75661	AGCCACCAAG	CCTGGCCTAA	GTGACATGTT	CTTATATTGT	TCCTTTCTTT	CALLACTURE
75721	CGACTGAGTC	TCACCCTGTT	GCACAGGCTG	GAGTGCAGTG	GCGTCATTTC	GGCTCATTGC
75781	AACCTCTGCT	TCCCGGGTTC	AAGCGATTCC	CTTGCCTCAG	CCTCCTGAGT	GCCÁCCACCC
75841	CCAGCTAATT	TTTGTACTTT	TAGTAGAGAT	GGTGTTTCAC	CATGTCGGCT	AGGCTGATCT
75901	CAAACTCCTG	GCCTCAGGTG	ATCCGCCCCC	GAGTCTCCCA	AAGTGCTAGG	ATTACAGGCG
75961	TGGGCCACGG	GGCCCAGCCT	TATATTATTT	CTTTTACTAC	AATATATTAG	TATGATGCAG
76021	GTGCTTCAAT	TGTTTATACA	CTTTCCATAA	TTTTGTATAA	TTCTTATACC	CTGTCACTCT
76081	GAGGAATAGC	CGGTCTAAGT	GTTTTTCCAC	CACTGCTAAT	TCATCCATCA	CTAATCTCAT
76141	TAGACTGTTA	ATTCCCAGAG	GACATAAGCA	CACAAGCAGA	CAATGTTTAC	AAATGTTGGA
76201	CAAATGTTAT	TTAATAAAAC	AATGGGGTCA	CCCTTAGTCT	AAAAGATGTT	TCACTTTTCA
76261	TTTGTCATTG	AACTCTTATT	TGTAGGTTCC	CTTTTGACTT	TCCCACAATC	TAAGGCTGTT
76321	CTCTTTAACA	CATATTTTCA	TGAAAACATA	TATTTGAGCA	GAAATTGTTG	GGGAGTTGTA
76381	ATATTACCTT	TGTCCCTAAA	TATGAATCTA	TAATTATATC	AAATATATGG	GCAGACAATT
76441	TACTITGCCT	TTAATCTCAA	GAAAAAAATA	GCAATTACTT	GGGGTCGGAG	AGTAAAATAA
76501	GAAGTAGTGA	ACCTTAAAGT	AGCAAACTTT	AGAACAGAAT	AGTTTCAGAG	GGGATGAGAA
76561	GAGGTGATTT	TTCAGCTCAT	CAACAACAGA	TCTTATAATA	AATTACATGT	TCTGGTACTT
76621 76681	TTCTTGTCTT	TCTGTGTTAA	ATTTTGCTAT	TAAAAAAAT	AAATTTCAAA	TACATTGTTC
	ATCTTAAAAG	TCAAGAGTGT	GTTTTATTAA	AGTCAGTTGC	TTTATTTGCA	ACTCAAAAGA
76741 76801	TATATTTGAG	TTCCCAACTG	GAGATTGTCC	TATATGGTAA	CTTGCGTAAG	GTATGGTTAC
76861	TAAAAGTAAC	CTACAATTTT	CATGGGCTGA	AATTCATTTC	TATATTGCAG	CGTACAAAAA
76921	TARATARATA	AAAAATGCTT	GTTTTCTTTG	AAAACATATT	ATCTCAGTGC	CTCTAACTGC
76981	CHARICIATT	GGCTTTTTTG	CAGGCTTAAG	GGCTCTCCCT	TGTTCCTTTA	TGATCTCTAT
77041	DEDGCCCDDM	AGACCTCCTG	CCTTACACAA	CTCAGAGGGG	GACCTCAGAG	CTCTTTAAAA
77101	GAGGGATTTC	ATACTOGCCTG	TAGAGAAGTG	AAAAGGATGC	CCCACCCCCA	TCTATGAAAA
77161	CCCCGACCC	TROCKROCOM	TGTCTTCAAA	TCAAAGATTT	AAGTCTGTAG	CCCCCCACCA
77221	GCCGTGGAAT	CCTTCTCCC	CATGAACCCC	CTCCCATCCC	GCCCTAATTG	CTTTGGACTG
77281	GACCTGTGTT	ACTTCCCCA	GTCCACAGTT	CCTGTGCGAC	TGCACGAAGA	ATTCACAGAG
77341	TTTCGCTTTT	TTCTTCCTIG	TGAAGAAACA	CARTTATCAT	GAAAATTTAG	GTGGAAACCA
77401	CCTTCAGGGG	CAAAGGAGGG	ATAAGGGAAG	CATGTGCCCA	ACCACCCCTG	GGAAAAAGAA
77461	TGCCCCAGAC	TTCCTTCCC	AACAGGTAAT	TIATAAGAAA	AACAGAAAGT	GGTCTCTGAC
77521	TTTGTGGAAA	ADDITION	GTTGGGGGAA AGAGCATGAA	1 TGGGGACGC	CIGGACGCGT	TGTTTTTGTG
77581	AAACCCAAGT	GATTTGGTGC	GGGGAATTTT	A TA TERESTEE	CCOMMMM	TTTCCTGACC
77641	AACACAACTT	GGGAGCAGCG	CAGCGGCTCA	CYCLCACCAS WYTHT LITTIC	CCCTTTTGTG	AGGTGGAACA
77701	ACCAATCAGA	GCGCGCCTGC	GCTCTATATA	TACAGCCCCC	GCCAGGCGGG	CGACCAGAGC
			COLCINIAIN	THUMBUBBLE	CIGCCCAGGC	GCTGCTTCAT

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77761		CCACTTGTAC				
77821						CGGCCAAAAA
77881	GGCTGGGGGT	ACGCCTCGTA	AGGCGTCTGG	TCCCCCGGTG	TCAGAGCTCA	TCACCAAGGC
77941		TCTAAAGAGC				
78001	TGCCGCCGGC	TATGATGTGG	AGAAAAACAA	CAGCCGTATC	AAACTTGGTC	TCAAGAGCCT
78061	GGTGAGCAAG	GGCACTCTGG	TGCAAACGAA	AGGCACCGGT	GCTTCTGGCT	CCTTTAAACT
78121	CAACAAGAAG	GCAGCCTCCG	GGGAAGCCAA	GCCCAAGGTT	AAAAAGGCGG	GCGGAACCAA
78181	ACCTAAGAAG	CCAGTTGGGG	CAGCCAAGAA	GCCCAAGAAG	GCGGCTGGCG	GCGCAACTCC
78241		GCTAAGAAAA				
78301	CAAGAAAGTG	GCTAAGAGCC	CAAAGAAGGC	CAAGGTTGCG	AAGCCCAAGA	AAGCTGCCAA
78361	AAGTGCTGCT	AAGGCTGTGA	AGCCCAAGGC	CGCTAAGCCC	AAGGTTGTCA	AGCCTAAGAA
78421		AAGAAGAAAT				
78481	GAGCCACCAC	TGATCTCAAT	AAAAGAGCTG	GATAATTTCT	TTACTATCTG	CCTTTTCTTG
78541	TTCTGCCCTG	TTACTTAAGG	TTAGTCGTAT	GGGAGTTACT	GAGGTATCAG	ACGAATTGGG
78601	TGACGGGGTT	GGAGAGTGGC	CGTGGTGAGG	TTACAGCATT	TAAACCTTTA	TTGCGGCTTC
78661	TAGGTCCCTG	ACCGGAGGCT	TTTCTCGCTG	GCGGATGGTT	TTGGGATGGC	AGTCCCGCCC
78721	CAGGCCTGTG	AACGGCAGAA	AAGACCGCAA	AACAAGAGCC	AGTTTCTTAG	TCTAAAGGGA
78781	TGTCCGGATT	GGACTAAAAA	ATTTTCAAAA	GTCCCGCCCT	GCTCCCGGGT	TGGTCCGTTC
78841		TGACTTTCAT				
78901	TGTTTTTTGC	TTTACTGTGA	CTTAAAAGTT	TTGCCTCTTT	TCTCTTTATA	TTAATGTCTG
78961		CGCTTTCCAT				
79021		CCTGGGAGGA				
79081		TTCCTGTGGC				
79141		CTCCACCCAA				
79201		CGCTAAACTG				
79261		CAGCCATTTT				
79321		TAAAAATCGA				
79381		AACAGTATTT				
79441	ATTTCTAATA	AAGGCCTTCG	TTAATGGTTC	CCTCTGTTTG	ACATCCATGG	TGCTTCTGAA
79501		CTAGCGTCTT				
79561		TATGGGGACT				
79621	CCACAATGAT	TAATATAGTG	AGTTGATTTG	TTAGTGATAG	TGACCACGGA	TTCATCCCAA
79681	GAAAGAGAAA	GGGGAGGAG	GCAAGCAGAG	AGACAGGAAG	ACAGAGGCAG	GGAAGAAGGA
79741	GAAAACATTC	TCCCATGGTT	TAAGTAATTT	TGTGTTGTTA	ATTTTACATT	ACAACACGGT
79801	TTAACATGGT	GAACCCTCTA	TTTTGGTGTA	AGGTTTAACA	TATGGACATA	TTTTTCCCAA
79861	GACCATTTAT	GAACTTTCAT	TTCTGCTTCC	CCCTTCTTCC	TCCCGTGCCA	CCCTCCACGC
79921		TTTGGCTGTT				
79981	ACTGTCTTAG	GTTTCTCAGG	TTTCTATTTT	GTTCCTTTAG	TCATTCCCAC	AATTCTTAAG
80041	GTAGAATTGT	ATTGTTTTAA	ACATTGTGTT	GTGTGCTATC	CTCAATGCTG	AGATGATTAT
80101	GTGACAAATG	GCAAGTGTTC	AACTAATACC	TAAATCTGTA	GTATCTTATC	AAGCCTAATG
80161	CTACTTCACA	ATGCCTACTC	CATTCACCTC	ACTITATCTC	ATTACTGGCA	TTCTGTCATC
80221		ACAAGTAAAA				
80281	TATTTATATT	TATTTATTTA	TTTATGAGAC	GGAGTTTCCC	TCTGTCACCC	AGGCTGGAGT
80341	GCTGTGGCAC	GTTCTCGGCT	CACTGCAACC	TCCGCCTCAC	GGGTTCAAGC	GATTCTCCTG
80401	CCTCCGCCTC	CCGAGTAGCT	GAGATTACAG	GGGCCTGCCA	CCATGCCCGG	CTAATTTTTG
80461	TATTTTTAGT	AGAGACGGGG	TTTCACTAAG	TTGGCCAGGC	TGGTCTCGAA	CTCCTGACCT
80521	CAGGTTATCC	GCCCACCTCA	TCCTGCCAAA	GTGCTTAGAT	TACAGGCGTG	AACCACCGTT
80581	CACAGACTCA	AATCATTTTT	ATTACAGTAT	ATTGTTATAA	TTGTTGTTTT	ATTATCAGTT
80641	ATTGCTAATC	TCTTACAGTG	CCTGATTTAT	TTAAATTAAA	CATCATTGCC	ATGTGTATAT
80701	AGAAAAAAAC	AGTGTATATA	CGGTTCAGTA	CTATCTGTGG	TTTCAGGCAT	CCACTGGGGG
80761	TGCAGTTTAT	TAAACATGCA	TTTACATTAG	TCTCCCCTTT	GGGAGACTAA	TTAACTGAGA
80821	TGTTGTAACG	TGACTTTAAT	AGCAGATAGA	GCTAATTTTC	TCTCATTACT	CTTCTTTTTC
80881	AGAATTTTCC	TGGTTATTCC	ATTTTTTATT	TTTCCATATG	TATATTAAGA	TCTCTTCCAC
80941	CTCCTCCTGT	TTCTCCATCT	CAACATCAAA	CAATTAAAAA	DAAAAAAAA	GCTGGGCGCG

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			•			
81001					GCGGGTGGAT	
81061					CTCTACTAAA	
81121					TCGGGAGGCT	
81181					ACCTTGCACT	
81241					GCAGTGGCTC	
81301					GGTCAGGAGT	
81361					TAAATTAGCT	
81421					GGAGAATCAC	
81481	GAGGCAGAGG	TTGCAGTGAG	TTAAGATGAC	GCCACTGCAC	TCCATCTGGG	CGACAGAGCC
81541	AGACTCTCTC	TCAAAAAACT	AAATAAATAA	AAATAAAGTT	ATGGTACATT	GAACTTCTGT
81601	GTTCCTTTCT	CCCTTAGATA	CTTTCATGGC	TACCCATTTA	ATTGATGTTC	TTATCATCTC
81661	CAAGAGTTAG	TCAGGAGAGG	AATCAACCCA	AGCAAAAATA	GCTGATTTTC	TAATTTTCCT
81721	TCAATGCCCT	TTGGGGTCTT	AATCCATTTG	ATTTATGTAC	TTTCAATTAA	TCCTAACCTC
81781					CAAATGAAAC	
81841					TTGGCCTTTA	
81901					GTTTTATTCC	
81961					TTGGAAATTA	
82021					TTTTGAAACT	
82081					CTTGCAAATT	
82141					TTAGTATATT	
82201					TTAGACTCAC	
82261					TGACGAATTT	
82321					AAAATGAAAG	
82381					ACCTTCCATA	
82441					TTTTACAAGG	
82501					GACCTCTTCC	
82561					AAGTAGAATA	
82621					TTAGTAATTA	
82681					TATATATT	
82741					GCATAAACAA	
82801					TATGAGTAAC	
82861					TATTTATTTA	
82921					ATGGCGTGAT	
82981					CAGCCTCCTG	
83041					ATTTTTAGTA	
83101	TTCCCCATCT	TTGCCAGGCT	ACTOTOCO	ATTITITIOT TO THE PROPERTY OF	AGTGATCCAC	GAGACGGGGT
83161						
83221					CGGCGCATTA ATTATACACT	
83281	GCTCTCGATA	TOCIAICAIG	GCIACAAATT	GAAGTATCAT	ATTATACACT	CCTAGGCAAA
83341					GTAAGGACAT	
83401					TACATCACTA	
83461	OUTGONGWIN	GGGATTIAGG	GTACAGTGGC	AACAACAGTT	TTGGGAACTA	GCATTTTTTG
83521					TCTATATTTA	
83581	CALICITGIC	ACAGCACTTT	GAAGTAAGTG	CCATTGTCAT	TCCCACTTCA	GGGTGAAGGA
83641	CHARAGCTTG	GIGICATIAA	GGATGTAGCT	AGTTAGCTGT	GTGTGTGTGT	GTGTGTGTGT
83701	GIGCATITIT	TTTTAAATTT	AAAGTCAATA	AATTTTTATT	TGAAGAATTT	CACATCAAGG
	CHARACTTTGT	TCCTCTAAAG	AGCTGGAGTC	AAAATGTATC	TTCAAAAGAT	TCATCTTCAA
83761 83821	GTTAGCCCTT	CITAATAGAA	CTGATGCTTA	ATCCACAGTT	GTCAGCCCAC	AGTTCTTTTA
83881	CCCCACCCC	TITITITIT	TTTTTTTGAG	ACGGAGTCTC	TCACTGTCAC	CCAGGCTGCT
83941	GGGCAGTGGC	GTGATCTCGG	CTCGCTGCAA	CCTCTGCCTC	CCGGGTTCAA	GTGATTCTCC
	TOTATION	TCCTTAGTAG	CIGGGACCAC	AGGCGCATGC	CATCGTGCTC	GGCTAATTTT
84001	TGTATTTTA	TTAGAGACAG	GGTTTCACTA	TGTTGGCCAG	GCTGATCTCA	AACTCCTGAC
84061	CTCATGATCC	GCCTGCCTTG	GCCTCTCAAA	GTGCTGGGAT	TACAGGTGTG	AGCCACTGCA
84121	CCCGGCCTTA	TTTTGCCTTC	TTTAATCTCC	ATTTGAACAT	ACACATACTG	ATGAAAACTA
84181	CAACATTCTT	CACCAAAAAT	CTTTGGGATT	TAATTTCTTC	AACCACTTTA	CTTTGGGGTC

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84241	ATTTTAAGAT	TAGGTGTATC	TGCCTGGTTC	TCAATTTGAC	ACCCTTTCTC	TCTAAACATG
84301	AATGAGTTCC	AATCATATTT	ATTCCTAAGC	TATCACACTC	AAATATACTA	CAGATCTGTG
84361	GAATATGCCA	AAAGTTAAGG	TGAAAAATTA	AATTATTAGG	TATTTCATAG	TTTTGCTAGT
84421	TTTTGATCTG	TGAGTGAATA	TAACTATCCT	CTATGTCCTG	GCACTGTTCC	TCAGAAACAT
84481	AGGGTCCACA	TATGTAATTT	TAAATTTTTT	AATAGGCACA	TTTTAAAAAG	TGAAAAAAGA
84541	AATCTATTTT	AATGATTTGA	ATCCAGTGTA	ACCAAAAATT	GTTTCAACAA	GGTATCTAAT
84601	ATTAAAATAT	TGAGTTTTTA	CTTTGTTATT	TTACTAGTTC	TTTGAAATCT	GGTGTGTATT
84661	TTACACTTAA	AGCACATCAC	AGTTTGGAGT	AGCCACATTT	CCAATGCTTA	ATACTCACAT
84721	ATGGTTAGTG	GCAACTATCT	TGGACAGGAC	AGCTTTTATA	CTCTGGGAAG	ACACAAGCAA
84781	ATACTTGCTC	TGCAGCAGAA	TCCAGATGTT	TTCCAAGAAA	ACACTTTTTC	TGACCTGTTC
84841	CTGAAACCCA	GGTAGTGTCT	CTAATACTTT	ATATTTTATT	GGTTTGTCCT	ATTGTAACCA
84901	CCCAACGGGC	TCTCCTTGTC	CACTTCCTAG	ACAGAGCTGA	TTTATCAAGA	CAGGGGAATT
84961	GCAATAAGGA	GCCAGCGCTA	CAGGAGACTA	GAGTTTTATT	ATTACTCAAA	TCAGTCTCCT
85021	TGAGAATTTG	GGGACCAAAG	TTTTTAAGGA	TAATTTGATT	GTAGGGGACC	AGTGAGTCGG
85081	GAGTGCTGCT	TGGTTGGGTC	AGAGATGAAA	TTATAGGGAG	CCTAAGCTGT	CCTCTTGTGC
85141	TAAATCAGTT	CCTGGGAGTG	GTGGGGTGGG	GGACTCAAGA	CCAGATAATC	CAGTTTATCT
85201	ATATGGGTGG	TGCCAGCTAA	TCCATTGTGT	TCAGGGTCTG	CAAAATAGCT	CAAGCATTGA
85261	TCTTAGGTTT	TAAAATAGTG	ATTTTATCCC	CAGGAGCAAT	TTGAGGTTTA	GAATCTTGTA
85321	GCTTCCAGCT	GCATGACTCC	TAAACCATAA	TTTATAATCT	TGTGGCTAAT	TTGTTAGTCC
85381	TGCAAAAGCA	GTCTGGTCCC	CAGGCAGGAA	AGGGGTTTGT	TTCTGAAAGG	GCTGTTATTG
85441	TTTTTGTTTA	AAAGCAAAAG	TATAAACTAA	GCTCCTCCCA	AAGTTAGTTA	ATCCCAAACT
85501	CAGGAATGAA	AAGGACAGCT	TGGAGTTTAG	ACGTTAGATG	GAGTCGGTTA	GGTAAGATCT
85561	CTTTCACTGT	AATAATTTTC	TCAGTTATGA	TTTTTGCAAA	GGCAGTTTCA	CTGTCCACTT
85621	CACCTCACAT	CAGGCCTCTG	ACTAGAGGAT	TCCAACAATA	CTTAGGCCAG	GACACCACCA
85681	TGTCTCCTTA	TCCACCCTGA	GGGAGTCCAA	TTTCTGAAAC	AAAGGAAACT	ATATATGATA
85741	GTATGAAACT	ATATATGAGA	AGGAAATTAT	ATATGATAAT	CAATTTTAGG	GTTATCTTAT
85801	TGATTAGAAG	ATATTAAAGT	GTGACACTGC	CTGGCAATGA	TATCTGCTGG	TAGTAAGAAT
85861	TTGGCGAATT	TAGTGAAATT	CCTGAGGCTG	AACCTCCACT	TCTGTAAAAT	GGAGACAGTG
85921	AGATAATTTG	CCTTACAATG	CTGAAGTAAG	AATTTTACAC	AATAATTCAG	ACCAACCACT
85981	TCATGTGGTA	CTTGGCCCGT	GGAAGACTAT	CAATGACAGT	TAGTTTATAG	TTTATACTAT
86041	TAATGAATCC	TTTGTTTCAT	TGTTATTTCC	TTCTACACGT	TGGCCTCTCT	AAAAGAAGGT
86101	AATATTCAAT	ACAAATAAAG	TTAAAACAGC	TTGCAGAGTT	GTCCCAGGGA	ACTCACTTAA
86161	CCACTGAAGT	GTTCAAATTG	CTTAAGGTTG	ACTTTATATT	CTCCTGACTA	ACCTTTCTCC
86221	TTCTGGTATT	TCTTCTGAGA	ACAGCACCAC	CATCCAAAGC	ATCATGCAAA	CAGTGGTCAT
86281	CCCAGACCAG	TAATTCTCAA	CTCACAGGGT	GCTCCTGCAG	AGATGTATTT	GAATAGAGTG
86341	GTAGGATGCT	GAAGAAGGCC	ACGTAAAATT	TGGCCAGTGA	TCTGGGGCAG	ATTTATCCTG
86401	AAGCTAATGA	AACACAAGTG	TAAGGGCCTG	TACTTCCAAG	GTGCAGAGAG	GGGCCCTACA
86461	AATGTGTTAG	TTTGTCTCTC	TCTCTCTCTC	TGATTTTAAA	ATTTGCAGTA	TTAAGGTACT
86521	TTAATCACGG	ATGGTTCAGG	CTGCTATTTT	CACTCAATCC	TCCTTTTTAT	TAAAATCACC
86581	ATTGTCTGAT	TATGTTAGAA	TCCTGATGAA	AATATTTGGA	ATTTGAGTAA	GAGAAAGTTT
86641	AGTTGAAGAT	GTATCTAGTA	TGGGGATAAT	AAGTTACGTG	ATTTGCATAT	GTGATCATGT
86701	GTACTTCATT	CGTTGCCAGC	CAATCTGACG	TAAGAATGGC	TTCAAGGAGG	CCGGGCGCGG
86761	TGGCTCACGC	CTGTAATCCT	AGCACTTTGG	GAGGCCGAGA	CGGGCGGATC	ACGAGGTCAG
86821	GAGATCGAGA	CCATCTTGGC	TAACACGGTG	AAACCCCGTT	TCTACTAAAA	ATACAAAAA
86881	TTAGCCGGGC	GTGTTGGCGG	GCGCCTGTAG	TCCCAGCTAC	TTGGGAGGCT	GAGGCAGGAG
86941	AATGGCATGA	ACCTGGGAGG	CGGAGCTTGC	AGTGAGCCGA	GATTGCGCCA	CTGCACTCCA
87001	ACCTGGGAGA	CACAGCGAGA	CTCCGTCTCA	AAAAAAAAA	AAAAAGAATG	GCTTCAAGGA
87061	ATGTTCCTAC	TGCTCACTGG	AATAACTCAC	CTAAATTCCT	GGCAAGATGC	AGGTCTAGAT
87121	AAAATGTTAT	GACATCTAAG	TATTCAAAAC	ACATTCCCAG	CACTGAGAGT	GAGTGTCTAG
87181	TGGAGAGTAG	AAACGTATAG	AGCCAGAAGC	TAGTCTGGAA	AGAATTCTTA	CAAAGTTTAC
87241	AACTTACATG	TGAAAGGAGC	TTAACAGAGG	ATTTTCCAAA	TTTGAAAACA	ATCCTAAAAA
87301	CTTACTTGAC	ATTACCAATA	ATGTGTTTTG	AAACTGAAAT	ACTTCTAAGT	TATGAAGAAA
87361	ACATATTATC	ATCAGCCACC	CTGGAGGAAA	GATTGAATTC	TATTTCCATT	ACCTATAGAC
87421	AACATTACAA	AATAATTTCG	ATCTGAAGAT	GGAATCAGAG	TATTCAGTCA	AAACTACAGG

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87481	AAAATATACT	TGGTAGTGT	: ATATTCAGAA	GTTAATAAA	TATGCTATTT	TCTGAATTTT
87541	GTGATGGCTG	TTGTTTTGT	AGCTTTTATA	AAATTGGAAT	TTGATTTTAT	TTTCCCATTA
87601	TAAATTTATA	TTTACAGTCT	GCAGTACTTT	TGCATTTTT	ATTTTACATT	ATAGTTTTTA
87661	ATAGTTAACA	AGTTGTAAAA	GGTTTGATCC	CCAGAAAACC	TTGATCTACC	CCATCAGTTA
87721	AGTATACTAA	TATATTTAGA	AAATGGATGA	AATCAGCATT	TGAATATTTT	TAAATATTTA
87781	TTAAAAGAGG	ACATGGGTAA	AAGAGCTTTG	CAGTTGCCAC	רכישיראיייריז מיייראיייריז	CAAATTCCCT
87841	GGATAAGGAT	GACCGCATAA	TCTTTGGATG	GTCATACGCA	AGTCTTGTGT	ACTTGTTACA
87901	TAAATCTATT	TAGTGGACTI	TTGGCAGTGT	GTACTGAGGC	CAGTTTCTTC	CACCTGAGCT
87961	CTGACTCCAC	CTCCAGCAGC	CCAAAACCAA	TACTGAATTT	TGGGGTCAGC	TATTGTTTTT
88021	GTGGACTTAG	GTAACTACAC	ACACATTGTC	TTTATGATAG	יי מאיים מיידים	ACTGCCATCA
88081	GAACTAAAAT	TGTCACGTGG	ATTAAAAGGA	GTGACGGTGG	TGTCCCCAGG	AGCCTTTCAA
88141	TATGTAAGTA	TTTACACATA	TACATGCTAA	AAAGACCCCT	' ACCARTOC	TAACAAGGGC
88201	AAAACAGTAA	CTCAGCTTGT	TTTCTCGCAG	TARARCCGGT	TGAAAAGGCC	TGATAGACTT
88261	GTCTGCAGTT	ACAAAACTTG	TGTGTAGTTA	TCACCTTTAT	CONTROL I	AACTAACATA
88321	GACAACCGAA	TGGGTTACAA	CTGTTTTTAA	GTGAAATTGT	RICICCION CACTCCCTCT	GAAAAGAGCC
88381	TTTTCAATGA	GGAAGAAACG	GGCAGACTTA	TGCCCTTTCC	CCACCCATCC	GACGTGCCAG
88441	CTGGATATCT	TTGGGCATGA	TGGTGACGCG	TTTAGCGTGA	ATACCCCACA	GACGIGCCAG
88501	TTCGAAGAGT	CCCACCAGGT	AGGCCTCACA	ACCUTCUTCO	ATAGCGCACA	CCGCAGAGCT
88561	CTGGAAACGC	AGGTCGGTTT	TCAACTCCTC	CCCCATTTCC	CCCACCACCA	GCTGGAACGG
88621	CAGCTTCCGG	ATCAGCAGCT	CCCTCCACTT	CTCCTACCCA	CGCACCAGGC	GCTGGAACGG
88681	GGTGCCCGGG	CGGTAGCGAT	CACCAMACLI	CIGGIAGCGA	CEGATTTCGC	CGCTCTTACG
88741	GGCTGCTTTA	GTAGCAAGCT	GAGGIIICII	ACCEPTECCC	GIGGCCGGAG	TGCGAGCTGT
88801	TIGCTTCGTA	CGAGCCATTT	GCIIGCGCGG	CACACACAAA	ACTOTACE	ACTGAGAGCA
88861	AGTGGCCTTT	AAATATAGTG	ACAAACAGAG	TCATTCCTCC	AGIGIAGIGA	CAAAAGTCCC
88921	GCGCGATAAA	ATCATTGGCT	CANCACACIC	CACACTCATT	COMMONMEN	TAGACAATCT
88981	TATTGGATGA	GTTGCCCCAC	CCCCCATCCT	CAGACIGATI	GGITCATTAC	CTGCAGCGAC
89041	AAATTGTCTA	AAATTOTAGT	TCATCCACTC	CCARACARCA	TITCAGTTAT	CIGCAGCGAC
89101	AAGGATTTTT	AAAATCTAAA	TTCCCATTC	CCAAAGAACA	GAGTGTATAA	AAATTCTGCA
89161	TTCCTGACAG	TOTOGONACT	TATCARTICA	GIAAGIIIGA	GIGGGACTIG	CCAGAAACGT
89221	TCAGACTCAT	GTCGGGAAAT	AACCOTTATA	UTTORCACACI	CACTAGACCA	CCAGAAACGT GCTATTTTGT
89281	TACTGGCGAA	CACCAACTTT	*COTTCCCCTT	TICAGAGAAT	GAGATTCCAT	CATTCCCACC
89341	CTGCCTGTTC	TCAABATGTC	TTATTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	TCCCCTTTAA	GTCCAAGTCA	TATACTCTAA
89401	AATGTACTTT	CTABAGGAAG	CTCTTATTIGGI	CTCCAAAC	AACTTTTTAA	TATACTCTAA
89461	CTAGGGGGGC	GGTGGCTCAC	GCCTCTA ATC	CCACCAMACII	GGGAGGGCGA	CACCATTAGG
89521	TCACTAGAGG	CCAGGAGTTC	DAGACDACCC	TCCCTARA	GGTGAAACCC	GATGGGACGA
89581	AAAAATACAA	AAACTAGCTG	GGCGCGCTTC	CACACCCCC	TAATCCCAAG	CGTCTCGCAT
89641	GCTGAGGCAT	GAGAACCGCG	TGNACCCCCC	CCCTCCTCCT	TGCAGTAAGC	TACACAGGAG
89701	CCGCTGCACT	CCAGCCTGGG	TCACACACA	A CA CECECECE	AAAACAAACC	CGATATCGCG
89761	AAAAGCAAAA	AATACCCTAA	CAGAAGCAACI	TTATCATCAT	TTCTTGTGTA	AATCCAAACG
89821	GCTCTGAAAA	ATGCCGTTTC	AAGTGTAAGC	TATCATCCT	GATTTGAGTG	ACTATGGACG
89881	CTTGGCCTTA	TCGTGGCTCT	CONTRACTOR	CARCACCACC	GCCTGAATAT	TTTACTTGAC
89941	GCCTCCCTGA	GCAATAGTGA	CGTTGCCCAG	CTCCTTCTTC	ACCTCCTCGT	TGGACAGGAC
90001	GGCCAGCTGC	AGGTGGCGGG	CGTTGCCCAG	CCCCCTCTTC	TCACGTATGG	CGTTTCGGAT
90061	CAGTTCTAAG	ATCTCGGCGG	CCACCTATTC	TARCTACACT	GGCGCACCGG	CGCTGCCCAC
90121	CTCAAAATAA	TTGCCCTTTC	COADDIATE	TANGIACACI	CCCTATTGGG	CTCCGACCGG
90181	CCGGTAGCGA	CGAACAAGTT	TTTTCCTTTTTT	CTCCATTTTT	CACGTCCGCA	AACTGCAAGC
90241	TATGAAAGCA	GCGGAAAACT	CTCAAACACA	PCC PCCMITTIC	AATGGCGCCT	AATAGCGACC
90301	TTTTATACAA	ACTGCAAGGC	TECANTAGE	ACCENTION	TTGGTCAATT	GAACAAATCC
90361	CTTTATCCAA	TAGAAAAACA	TARCATARA	TCCMTATTCCTA	CATAAACCCC	ATGTTTGGTG
90421	GAAACCGTGT	TTCTTTTCTC	CAATCAGAAG	TCACCAATC	TANACCGTCA	ACCUCTCAGT
90481	AGGACTATAA	ATACATGGGC	TCTCAGAAG	TOWNSHAT CT	ACTCTGTAGT	COLCACTOR
90541	AGTAGCTTTT	CTATTCTGTT	TAGGARCIGI	DECCORORS	CCCTCTAAGT	COCCOCCACC
90601	CCCTAAAAAG	GGTTCTAACA	AGGCTATCAC	TARGCCIGAA	AAGAAGGATG	CIGCICCAGC
90661	TAAGCGCAGC	CGCAAGGAGA	COLTAICNE	CTATCTCTA	AAGGATCTGA	A CONCOMOCS
-			CIMITOTAL	CIMIGIGIAC	MAGGITCIGA	AGCAGGTCCA

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90721		GGCATCTCAT				
90781						GCTCGACCAT
90841						TGGCTAAGCA
90901		GAGGGCACTA				
90961		CCAAACCCAA				
91021						AGAGGATCAA
91081		AGCGAAGACA				
91141						CACGAGGTCA
91201	GGAGATGGAG	ACCATCCTGC	TTAACACGAT	GAAACCCCGT	CTCTACTAAA	AATACAAAAT
91261	AATTAGCTGG	GCGTGATGGT	GGGCGCCTGT	AGTCCCAGCT	ACTCGGGAGG	CTGAGGCAGG
91321	AGAATGGCGT	GAACGCGGGA	GGCGGAGCTT	GCAGTGAGCC	GAGATCGCGC	CATGGCACTC
91381						AAAAAAAAA
91441		TTGAAGCAGA				
91501		TCCTTTGTTA				
91561	GGTTTCTGTA	CTATAGTCCC	TTCTGTGGTG	GCCAGAAATA	TGTTACAGGA	AAGAGGTCCC
91621	CATCCAGACC	CCAAGAGAGG	GTTCTTGGAT	CCCGCGCAAG	AAAGAGTTCA	GGGTGAGTCC
91681	GCAGTGCAAA	GTAAATGCAA	GTTTACTAAG	AAAGTAAAGT	GGTGAAACGA	CAACTACTCC
91741		CAGGACATTC				
91801	ACTTGTATAT	ATGGGGAGAT	GTGCTCTGCT	ACAAGTTTGT	GATAAAGGAT	TAATTTTCTT
91861	AGTTACTATA	TTTTGCAAGA	ATCAACATTA	TTATCTTTAA	ACAAAATTAA	GAATGCCTTT
91921	GTTCTCCAGA	TATAGGGATA	TCTGGACACT	CCTAAGTCTG	AGTCTGTTTA	GTAAACATTA
91981		CCCTTAACCG				
92041	GCAGCCCAGA	AAGTCTCAGC	CTCATTTTCC	TAGCCCTCAC	TCAAAATGGA	GTTACTCTGG
92101		TCTGACACTT				
92161		TTTTGAAATA				
92221		AAGCCAAAAC				
92281		GATACTCGGC				
92341		TTATTACCAT				
92401		CAAAACTTGC				
92461		GTCCCATGTA				
92521		TTTTTTTGTT				
92581	CAATGGCGCG	ATCTCGGCTC	ACTGCAACCT	CCGCTTCCCA	GGTTCAAGCG	ATTCTACTGC
92641		CGAGTAGCTG				
92701	ATTTTTAGTA	GAGACAGGGT	TTCACCGTGT	TGGCCAGGAT	GTTCTCAATC	TCCTTACCTC
92761	GTGATCCGCC	CGCCTCGTCC	TGCCAAAGTG	CTCGGATTAC	AGACGTGAGC	CACTGCACCC
92821	GACCAATCTG	TCTTTTTGTA	GAGGGGCCTC	AAGCATGAAC	TTACTGATGG	GTGAGAAAA
92881	CAGAATTTTC	TTTTCCCCTA	CAATATAAAC	ATTAATTGTA	ATGTTATCAT	TCAGGACATT
92941	TTGGTGACCA	ATCTTACAGA	AATTTTATCT	TGTGCAAGTC	TATGCAAACC	AATATGTAAA
93001		GTGAGATTGT				
93061	TTCTAATGAT	TATTTTCATT	ACTGCATTTC	ATTGTAGGGA	AGTAGATAAT	TGCCCTTTAT
93121	TCACTGACCT	TCGCTTTTTA	AAATTTAAA	CCATGTTACC	ATGAAAATGC	TTTTCAGTAT
93181	TTCTCTACAC	ACAAGATTGC	TGTAAGGGCA	AAAATAGAGA	TAGGAATCAT	GCATCCATTG
93241	ATATACATAT	TTTGATTTTT	AATACATGTT	ACCAAGTTGC	CTCCTGAAGG	TCTGTTTACA
93301	CTCTCACCAA	CAGGGTGTTT	TTTCCTGACT	TCCACAAATG	CTCTTGAACA	GTGGGTGTGT
93361	TAGTCTGTTC	AAATTGCCGA	CATGAACAAT	TAAATCTCAT	TGTTGTTTTT	ATTTTTAAGA
93421	CAATTATTGT	TTGAGACTGC	ACATTTTGAT	AATAACATTT	CTTCTATTAT	GGTTTGATTA
93481	CTCATGATTC	TTGCCCATTT	TCTTTTGGGA	TGTTGCCTTA	TGTACATTAT	TTTAAATAGA
93541	TAGCTCCATG	TATTAAAAGA	TTATTAAGTT	TGAGGGCTTA	TGATATGTCA	GTTACATTTC
93601	TAAGATTTTT	TTTTTTTTT	TTTTTGAGAC	GGAGTTTCAC	ACTTGTTGCC	CAGGCTGGAG
93661	TGCAATGGTG	CGATCTCGGC	TCACCGCAAC	CTCCGCCTCC	AGGGTTCAAG	CAATTCTCCT
93721	GCCTCAGCCT	CCCCAGTAAT	TGGGACTACT	GGCAAGCGCC	ACCACGCCTG	GCTAATTTTG
93781	TATTTTTATT	AGAGATGAGG	TTTCTCCATG	TTGGTCAGAC	TGGTCTCGAA	CTGCCGACCT
93841	CAGGTGATCC	ACCCGCCTCG	GCCTCCCAAA	GTGCTGGGAT	TACAGGTATG	AGCCACTGGG
93901	CCCGGCCACA	TTTCTAAATT	CTTTATAAGT	ATAAATTCAT	TCAATCTTCA	CCAAAACTCA

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93961	ATGAAGTGTG	AGTACTATTA	TTATCATTGT	TTTACAGATC	AAAACAAGTA	ATACAGTCAC
94021	TTACTGAGTT	CTATACACCT	GGTAATTTTT	TTGTTTCGTT	GTTCTATCAA	TTATTGGGGA
94081	AGGGGTGTTG	AAATCTCTAC	CTTTAAATCA	TGTATGTGTC	TATTTCTCCT	TTCGGTTCTA
94141	TCAGGTTTTG	CTACACATAT	TTTGCAGTTC	TGTTATTTGG	TGCATATACA	TTTAGAATTG
94201	CTTGTTTTTC	GTATTGGATT	GACCCTGTTA	TCATTATGTA	ATATCCCTGT	CTGTTCCTAG
94261	TAATTTTCTT	TGCTCTGAAA	TATACTTATC	TGATATATCA	TCCAAAAGAC	CACCAGGATG
94321	GCTAAAGAGT	AGAAAGGAGA	GATTTACTGG	CAATACTAAT	TTGCAAGCCA	GGAAGAGATG
94381	GTCCCAGAAC	CTGCCAAAAT	TACTCTCTCT	TTGGGGAGAA	GGAGCAGGTT	GGTTATTTTT
94441	ATGCCTCATA	GGCTATATAT	TACACAATAG	AGTCATACAT	ATTTAGCACG	TTTGGGGGGA
94501	CAGCTATATA	TATTATGAGG	GGTGCCAAGT	GCATTCACAA	TGGATAAACA	CGTGTAATAT
94561	ACCTCCCATG	TTCACTTCGA	GGTTAAATTT	TGGTTAAAAT	GAGGTAGAAT	TTAGGTCTTT
94621	ACATCACAAG	GTGAACTATA	GGAACAAAGT	TTACGTGCTG	CCTCTAGCAG	CTGGCTGAAA
94681	ATGGCTTAAG	GTCTACAATT	ACGTGTAAGA	ATAGAATGTG	TGTCAAGGCG	CIGGCIGAAA
94741	CAATCAGAGT	TGTAGTGGAC	TGGACTGTAA	ATCAGAGTTA	GGAGGGGCTTC	TCATACCTCC
94801	TATAGTTAAG	GAATTTAGCA	AGTGTGAGTT	TTTTGGTAGT	CTTTCCAATT	TAGGAAMTTG
94861	CCATGCCAGC	CAAGCCATGA	ATGCTCTACC	AGTAGGTAAC	TTTCTTCCT	TAGGAMITIG
94921	GTCTGTCTTA	GTTGGTATAG	GGGCATCTAT	TTTGGTCTTT	CAGATCCCAG	TAMICITAGA
94981	TACAGATACT	CTTGCAGTTT	TGGGCTGATG	TTTATATGGC	THE TOTAL	
95041				AGTGAGATTC		TGCAGCCTTT
95101	TTGTTTTTT	TTTTTTCACA	TECANTOR	CTCTTGTTGT	CCACAGACAG	TGTACAGTTG
95161	ACAGTCTCAG	CTCACTCCAA	COTCCCCTC	CTGGGTTCAA	CCAGGCTGGG	GTGCAGTGGC
95221	TCTTGAGCAG	CTCGCGATTCC	ACCCATCCCC	CACCACACCC	GGGATTCTCC	TGCCTCAGCC
95281				CTGGTCTCGA		
95341				TTACAGGTGT		
95401	ACTGTTTTTT	TATEGETETA	TTTTATACCAC	ACACATTTAA	GAGACCTCGC	GCCCAGCCAA
95461	GGCTTAAGTT	CATGAAGGGT	ACTOTOCOA	CCATAGTCTC	TGCAATTATT	GATATCTTAG
95521	CAGAAATCAC	TGACAAGGCA	CATTOTOGAA	TAGGTGAAAA	CCCACT	AAATGTTTGC
95581				TTACCTAACT		
95641	TTATATACCA	THOUGHOUND TO	I CAGAAI IAA	TIGGGGCTTA	TCCCAATGAG	TTATAGATGC
95701	TGGGAGGCAA	ANGREGATE	CCTTCCAAAGAA	TIGGGGCTTA	GATTCTGGTA	AAACAGGTTA
95761	CAGAAAGAAC	AAGAGGIIIG	GC11GCAAAG	GTGGCCTTGT ATGATTTTTA	TAGGTAGGTG	AAGCCTCCCT
95821	TCCTGGATCT	GGGGAAAGCT	ATACARACCE	GAGGAGGCAT	AGTGTCAGAC	TCTCAGTCTC
95881	TCTACAGATG	TOUGHAMAGGI	ATAGAAAGGI	GCAGCTTTGC	GGCTGCATTA	ATGGAGATTC
95941	GCCAAGCAGC	ACCCATTO	A A A TO	AAGAAATATA	AAGCCCATTT	CTGCCTGCTG
96001	ATTTCCTTTA	CACTECTECE	AAAIAIGICA	AAGAAATATA	TTTTGGGGTA	AAATATTTTG
96061	CCCTTCCTTA	CTC3 A C B TCT	CITATAAGAA	AAGGAAGAGA	CACCIGAGCT	GACACACATA
96121	TGCCCTTTAGC	TTCCCACCATGI	COLOR	TAAGAAGGCC	CTCACCAGAT	ACTAATTCCA
96181	CCCACTCTCT	CCTATTO	CTAGAACAGT	AGGAAATAAA	TTTCTTTCT	TTAAAAGTTA
96241	ATCTTACATC	ACTEATECE	TATAGTATCA	CAAAATGGAC	TAAGTAACTA	TATTATGATC
96301	TTAGAGGTTC	CTCTCCCCC	CCTACATCAT	ACACATACAC	AGGCCACATT	TGGAACATTG
96361	TGAGTATOTT	CARTACTATA	TACAAATGTA	CTACAAATTA	TATATGTATT	TTTAAATTTT
96421	TATTCAATAT	CCATASTTAT	TACTORON	CTTTTGTAGT	CAAAATGTCA	TTATAACATG
96481	TGGTTTGGAT	ATTTCTCCCC	TAGTCAGATG	TTTTACATTC	TTTCTTCATA	CTAAGTGATA
96541	TGAACCCTCC	TCARACCTET	TCTAAATCTC	ATGTTGAAAT	GTAATCTCCA	ATGTTGGAAG
96601	TGAAGCCTGG	ATCCCTTCT	TTGGATCGTG	AGGGTGAACC	CCTCATGAAG	CGCACTCTTC
96661	AGGGTAATCA	AIGGGITCIC	ACTTTGAGTT	CACAAGAGAT	CTGGTTCTTT	AAAAGAGTGT
96721	GACACCTCCC	CCATCTCTCT	CGCTCAGCTC	TCACCATATG	ATATGCCTAC	TCCCTCTTCA
96781	CTCCTCTACA	CCCTCCLCL	TTTCCTGAGG	ACTTGCCAGT	AGCAGATGCC	TGCACCACAC
96841	CTCCTGTACA	ATTOCCACAA	CCGTGAGCCA	AAAAAAATTA	CTTTTCTTTA	TAAATTAGTC
96901	AGTTTCAGGG	ATTCCCTTAT	AGTAATGCAA	GAACGAACTA	ACACACTAAG	TCTATTTCAT
96961	ATTTACAGAA	TAGUICAATC	TGAAGTACCC	TITTTCAACT	TCACAGTAGC	TACTTGTAGC
97021	TAGTGGGCAC	TGATTTGGAG	CGTGTTCAAG	GGTGAATTGT	ATTATGCAAT	TAACAGATTT
97021	TTTTTTTTTT	ATTCCCAAAC	CACGAGGCAT	AGATTGTCTT	ACTITCTCTG	CTCCTGGTGT
97141	TGGAGTTGTT	COMPOSES	AACTTATTTT	CCTCTTATAT	TTATATGGAA	TAAATAACCC
~ / ###	CCAATATTTC	CCTCCCCAAT	ATCTGCCTTT	TGTATGTTTT	TTGAAGGCAA	GTGCCTAGAA

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97201	TTTACTGTTT	TTGAAGCACT	TACTGAAAGG	ATTGCCATCA	AGTTGTTTTG	CTAATAGTAC
97261	ATGCCAGGCG	CTTGTTGGTT	TGCTTAATTC	AAGGTAACTT	GGATGAGAAG	AAGAGTTTTT
97321	CTCATCCATG	GCTCAGTGGA	GTATAGATTA	CTGATATTGT	GACTGGATGT	ACTCCTGCTT
97381	TCTAGTCTGA	GTTTTTGAAG	CTACCCTTAA	TCTTGGTTTC	AATTTTATCT	AGCCCTGTAC
97441	ATATCCAAGG	CTCTTTCCAA	AATGGTCTAC	GATTTGTTTA	GGAAGTTAGA	ATAGCTGTAC
97501				GACTTCAAAC		
97561				TCCTTGCCTT		
97621				TCTATTCTTT		
97681				TACATTTTCA		
97741				TAAGGAAACA		
97801				GCACTTAGCA		
97861				TCATGAACTC		
97921				GTATGATTTA		
97981				TCATGTACGT		
98041				CTTTTTAACA		-
98101				TGATGTTTAC		
98161				TATTGTGAAG		
98221				TAAAAATGAG		
98281				CAGCGTTGGC		
98341				GCCĆAGGCTG		
98401				AAGCGATTCT		
98461				CCAGCTAATT		
98521				TACGAACTCC		
98581				GGCATCAGCC		
98641		•		TCTGTCATCC		
98701				TCCTGGAATA	- · · ·	
98761				AAAGGAAAAA		
98821				AGAGCCAGAC		
98881				TCCCAAGGTT		
98941				ATTTATTTCA		
99001				ATTGCCCATT		
99061				GCTAACAGAC		
99121				TCAGTGCAGC		
99181	GGACAAGACT	CTCTTTTAGA	ATGGGGGGTC	TTATGACCTA	CAGGCAAACA	AGGTAGGTTA
99241				CTAGGGAAAA		
99301				GCTAGACTTT		
99361				CTTTTATAAT		
99421				TTCCTGAAAG		
99481				TCAAAAATGT		
99541				AAATTCAAGG		
99601				CAAAAAGGAA		
99661	TCCCTGGGGA	ATCTCATCAA	CCAGAGAAGA	TTAACTGTAT	CACAGGAGAG	GAGACTGGTG
99721				GCTGTCACCT		
99781				AAGTTGCCTA		
99841				AGTTCACTGG		
99901				GTCTTTGAGA		
99961				AACTCACTGC		
100021				AACTGGAACT		
100081	CCGGCTTTTT	TTTTTCTTTT	TCTCCCCCGT	TTCTTTTTTG	GTTATTTTAC	TGGAGACAGG
100141	GTTTCTCCAT	GTTGTCCACG	${\tt CTGGTCTCGA}$	ACGCCTGACC	CGCCGTCCTC	GGCCTCCCAA
100201				GCCCGATTTG		
L00261	TTTAGAAATT	GGTCGGAGTC	CACTCCTTTC	CAAAAACATG	AGTCACAATC	CGGGAAAAGC
100321	ACGAGCGGCT	GAAAGTCAAA	ATAACCAGAA	CAAAACCTCC	ACTCATGCTT	AAAAAAGGTA
100381				TATTAGTATT		

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100						
100441		GATTGGGTCA				
100501		GATTCTGATT				
100561		CTCCAGTCAG				
100621						GCGGCAAACA
100681		GCTCGCGCCA				
100741		GTGCACCGCC				
100801		TATCTCGCGG				
100861		GCCCGCGACA				
100921		GACGAGGAGC				
100981		AATATTCAGG				
101041						CAGACAAAAT
101101		GCAAAGGCTC				
101161	TGCTTTGGAT	TATGCCGCCC	ATAAAGATGT	TTTTGAGGTG	TTTTTAATGG	CTTTGAGTGT
101221	GGCACTTTTA	GTAATTTGTC	CTGCAGAAAT	TAGATCCATA	GAAACCTCAG	GAATTCTAGG
101281	TATGTGGGAG	AAGTGCCATG	CAGCACAAAA	CATGTTTACA	GGGGTGATTC	GCGTTAAGTT
101341						TGCATTCCTA
101401		ATGGAAGTGT				
101461		AGCAATTAAT				
101521	TTGAGTATGT	ATTTCCCAAA	ATGAGCTTTT	TTCCAGTTTG	GGGATGTTTT	GCTTTGTTTT
101581		CTCCCTCTCG				
101641		CGGGCTCAAG				
101701						TTATGCGACA
101761	CCCAGAAAAA	TACAATTTTA	AATAAAGCGC	ATATGCAAAT	TTCCCTAATC	GTCTCCAATA
101821		TCTTTTTTAT				
101881		AATTTTAAGA				
101941		TAAGGGGTTA				
102001	CTCTCTTAAT	CTGCAACCAG	GCACAGAGAT	GGACCAATCC	AAGAAGGGCG	CGGGGATTTT
102061		TGGGTCCAAT				
102121		CAGACGTCTC				
102181		CTCGGAAATC				
102241		AGAGCGCGCC				
102301		CTCTGCGCGA				
102361		TCCAGCGCCT				
102421		CTGCGGTGAT				
102481		CAAACCTTTG				
102541		CTCGCCGCAT				
102601		CCCAAAGGCT				
102661		TGTTGTCTTA				
102721		AAAGTTGATT				
102781		TATTAAGGTG				
102841		GCTATGATTA				
102901		TTAACTGGCG				
102961		GCGAGTTCCA				
103021	ATACATAGTC	CCCTAGGTTT	TCTCATATAT	TATATATATA	מדמדמדמדמ	TATATACTCT
103081	TAAATTCATT	TGGCTGTTAA	CATTAACCTG	חדיים מדיים ממ	TEGTECANA	TETERECER
103141	GGATCTAACT	GGCTCTCATT	TTATCCATAG	CTAGCTACCC	ACTTTAAATC	TGTCAGTCTG
103201	TCGACCAAGC	ATAATTTAAT	CCCTTATATA	TGAATTTTA	TATGTGTGGG	TOTORGICIO
103261	AATAGTCTAT	CTGGTTGCAT	TGCTTTGTCT	CCTCTAGGAC	TATGCACCAT	GACATGCCAC
103321	ATTCTTTTTT	TCAGTACTTC	TTGCCTGTAG	TTATTADAAT	CTAGAATTTA	CAAGTTTTAA
103381		TCTGTTGATC				
103441		AGGGATGGGA				
103501		TATTTTTTTA				
103561	AACAACCACA					
103621	TTTTTACAAT					

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103681				AACTGTCAAC		
103741				TAAATTCATT		
103801				CATTTTAAAA		
103861				CTTATACAAG		
103921				TAGAGATGCC		
103981				TGAAGGTCAA		TCCAGTTATA
104041	GATGTGCTGG	ATCTGATGTA	TAGCGCTTGA	CTTTTTATAT	TTTCTTTATC	TGTAGGAAAC
104101				TAGCATAAAA		
104161				TCAGTCCAGC		
104221				GGATTCTGCT		
104281				TCTCATTTGT		
104341				AATATTATAT		
104401				CAAAAAGGGG		
104461				ATTCTTTTGA		
104521				TTCTTTCTGA		
104581				GAAAAGACAT		
104641				GGAGTCATAG		
104701	AACAAACCTT	GTTAAAATAA	TCATATATTC	CTTTAATCTC	CTCATATGGT	TTACTTTCCC
104761	ACAATTGCCT	CTCTTTAACT	TAATGTGAAA	GCATTTAGCT	TTTGCCATTT	CTTTGGGGCT
104821	TCACTTTTTT	ATGAGGGTTC	TCCTGTCCCA	TAAAATTTAC	ATTAAATACA	TTTGTATGCT
104881	TTCATTCTGC	TAATCTGTTT	TATGGCAAAT	GAATTATCAG	GTCCAGCTGG	AGACCCTAAC
104941	AGAGTAGAGG	TAAAATTTTG	CCTCCCTACA	AGATAGAGAT	TGTGTGCATT	AAATGTTGTT
105001	TGTTCCCAGT	TGTTCAGTTT	GTCAGGCCTC	TGAGCCGAAG	CTAAGCCATC	ATATCCCCTG
105061	TGAACTGCAC	GTATGCCTCT	AGATGGCCTG	AAGTAACTGA	AGAAACACAA	AAGAAGTGAA
105121	AATGCCCTGT	TCCTGCCTTA	ACTGATGACA	TTACCTTGTG	AAATTCCTTC	TCCTGGCTCA
105181	TCCTGACTCA	AAAGCTCCCC	CACTGAGCAC	CTTGTGACCC	CCACCCCTGC	CAGCCAGAGA
105241	ACAACCCCCT	TTGACTGTAA	TTTTCCACTA	TCTACCCAAA	TCTTATAAAA	CGGACCCACC
105301	CCATCTCCCT	TCGCTGACTC	TTTTCGGACT	CAGCCCGCCT	GCACCCAGGT	AGAATAAACA
105361	GCCTTGTTGC	TCACACAAAC	CCTGTTTGAT	GGTCTCTTCA	CACGGACGCG	CCTGAAACAG
105421	TTTAACAGGG	TTTTTCCTGC	CCAGTCACAA	CAAAGTGATG	TTATGCTGCA	GGCTGAAGTT
105481	TACAGCTAAT	GCTGTTGAAG	TCTAAAATCA	GTTTTGGTTT	GTTAGATTTG	GGTGAGATGG
105541	CTAAGATTCT	CAGAGAAAGA	AGTCAAGTTT	GGGGTGCATT	TTTCAGACTT	AAAAATTTAG
105601	CAGTAGCCCT	TGCAGTTTTT	CCAATAGAAG	TGATTTAAGA	ATGTTTTCAG	GAAATTTAAA
105661	ACAACAGTGA	GAAGCGTGTA	TGGAGAGTTG	AACTACACTC	CAGACTTGGC	TATAGGAAAG
105721	CACGAATGCT	GCTATTGTAT	TGCACCTTGG	AAAAGAGAAC	AAAGGAATAT	TTTCGGACAA
105781	TTTTAACATG	TCACATATGA	AAAGCTAAAC	GGAATCTGTC	AACACCTTGT	ACGTTATTAC
105841	AGGCTGTGAT	TTTAAAAAAA	CAATCCTTAC	TAATACATAC	ATAGTTGCTG	CTAGCAATAT
105901	AGTGTTGGGA	GTAAAAACAC	GAAAATGAGA	GTTCAGGACA	ATATCCCAAC	TCTGAGCAGA
105961	TTTTTTTAAG	TAGTAACATC	TAAAATTAAA	CCATATTATG	TAATATTTAT	TTCTTTTCCA
106021				CTAATTAAAA		
106081				ATGAGCTATC		
106141				CCCATCCTCT		
106201				GGAGAAAGGT		
106261				GTAATGAAAG		
106321				ATGTAAAAAT		
106381				GCCCAGCAAG		
106441				TGAGAAGAAT		
106501				CTACTGCTTC		
106561				ACGCCTGCTC		
106621				AAAAAGTTTA		
106681				CTCAGTCGAG		
106741				AAAGGGATTA		
106801				CGAGGGGCGG		
106861	CGCCTCCTTA	TCCTCGCTCC	CGCTTTCAGT	TCTCAATAAG	GTCCGATGTT	CGTGTATAAA

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10	06921	TGCTCGTGGC	TTGCTTTCTT	TTCGCGTACC	TGGTTTTTGT	TGTCAGCTGG	TTAGACATGT	
10	06981	CTGGTCGCGG	CAAAGGCGGT	AAAGGTTTGG	GTAAGGGAGG	TGCCAAGCGT	CACCGAAAAG	
10	07041	TGCTGCGGGA	TAACATCCAA	GGCATCACCA	AACCGGCCAT	TCGGCGCCTT	GCTAGGCGTG	
10	07101	GTGGGGTTAA	GCGAATTTCC	GGTTTGATTT	ATGAGGAGAC	TCGTGGCGTT	CTCAAGGTGT	
10	07161	TTCTGGAGAA	CGTGATCCGG	GACGCCGTGA	CCTACACGGA	GCACGCCAAG	CGCAAGACTG	
	07221				AGCGTCAAGG			
	07281				ATGGCCCTTT			
	07341				ATGGTAACAT			
	07401	ATTCTGCCTA						
	07461	GAGTGGCTGC						
	07521				AGCAACTTAA			
	07581	AATGAGTAGC						
	07641				GGATTAAAAT			
	07701	CAGGCGTATC						
	07761				CAGAAGCCTT			
	07821				CCCTTCTCTG			
	07881				GAGGCTCCCC			
	07941				AGTCTGTTTT			
	08001				ATAAGATGGA			
	08061				GTCAACCCAA			
	08121				TGAAAGTTTC			
	08181				GTCAGCCAAA			
	08241	GTAACAAATA						
10	08301				TAGTGAATAG			
10	08361	GGGTGAGAGG	TCTGAGCTGG	AGATAAAAAT	GTGTGAGTCA	TCAGCAGATA	AATAAATGCT	
10	08421	GAGACCAGAT	GAGATGGCTA	AAAACTGAAA	CATAATGTAG	TGCAGCATTG	TTTGTAATAG	
10	08481	TAAATGAGTG	GCAACTGTAA	AGTTTTCATC	AGAAAGGACT	AGAGTGATCT	ATACATCCAT	
1.0	08541							
	70241	AAAATAGAGT	ATTTCTCTAC	ACAGCCCTAC	TAAAGAATGA	GAAAGCTGTA	CTCCACTACA	
	08601				TAAAGAATGA ACTCCTCTTT			
10			TACTCTGGCT	CAGTTCTTGG	ACTCCTCTTT	TCTTGGCTAA	CTCAACTGGC	
10 · 10	08601	TACTCTGGTG CTCACCACTT	TACTCTGGCT ACATGCTCTG	CAGTTCTTGG TGCTCTGTCA	ACTCCTCTTT	TCTTGGCTAA TCAACAGAAC	CTCAACTGGC ACCACGGCCT	
10 · 10	08601 08661 08721	TACTCTGGTG CTCACCACTT	TACTCTGGCT ACATGCTCTG GCCACGTTAA	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC	TCTTGGCTAA TCAACAGAAC TGTGATAGTG	CTCAACTGGC ACCACGGCCT GCAGCTTCGG	
10 · 10 10	08601 08661 08721	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC	
10 · 10 10	08601 08661 08721 08781	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC	<u> </u>
10 · 10 10	08601 08661 08721 08781	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC	
10 10 10	08601 08661 08721 08781 08 <u>841</u>	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CAAGGTGGCT	<u> </u>
10	08601 08661 08721 08781 08841	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG CACGCCTGTA	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG ATCCCGGCAC	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG TTTGGAAAGC	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT TGAGCCTGGT	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CAAGGTGGCT GGGTCAGGGG	
10	08601 08661 08721 08781 08841 08841	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG ATCCCGGCAC GCCTGGCCAA	CAGTTCTTGG TGCTCTAGCAA TGCCTAACCA AAACATAGAG TTTGGAAAGC TATTGTGAAA	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT TGAGCCTGGT CCCCGTCTCT	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAAA	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CAAGGTGGCT GGGTCAGGGG AAAAAAAAAA	
10	08601 08661 08721 08781 08841 08901 08961 09021	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTGGGCA	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG	CAGTTCTTGG TGCTCTAGCAA TGCCTAACCA AAACATAGAG TTTGGAAAGC TATTGTGAAA CGACTGTAAT	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG GGATCACCTG ACTAAAAAAA CGGGGAGGGTG	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CAAGGTGGCT  GGGTCAGGGG AAAAAAAAAT AGACAGGAGA	
10	08601 08661 08721 08781 08841 08841 08901 08961 09021	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTGGGCA ATAGCTTGAA	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAAA CGGGAGGGTG ATCGCGCTAT	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CAAGGTGGCT  GGGTCAGGGG AAAAAAAAT AGACAGGAGA TACACTTAGG	
10 10 10 10 10 10 10	08601 08661 08721 08781 08841 08841 08901 08961 09021 09081	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTGGGCA ATAGCTTGAA CCTGGGAGAC	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG ATCCCGGCAC GCCTGGCCAA TGGTTGCGG CTCGGGAGGC AAGAGTGAAA	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT	CTCAACTGGC ACCACGGCCT GCAGCTTCCAC CAAGGTGGCT  GGGTCAGGGG AAAAAAAAA AGACAGGAGA TACACTTAGG AAACCATCTC	
10	08601 08661 08721 08781 08841 08841 08901 08961 09021 09081 09141	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTTGGACA ATAGCTTGAA CCTGGGAACC CCTGGGAACC	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CAAGGTGGCT  GGGTCAGGGG AAAAAAAAT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA	
10 10 10 10 10 10 10 10 10	08601 08661 08721 08781 08841 08901 08961 09021 09081 09141 09201 09261	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACCTGTA CACGCCTGTA TTCGAGACCA TAGCTTGGACA CCTGGGAGAC CCTGACCTTA TGGGCATCTC	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC	CAGTTCTTGG TGCTCTACCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA AAAATATGTT	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT CACCAAGTCT	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CAAGGTGGCT  GGGTCAGGGG AAAAAAAAAT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC	
10 10 10 10 10 10 10 10 10 10	08601 08661 08721 08781 08841 08901 08961 09021 09081 09141 09201 09261 09321	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACCTGTA TTCGAGACCA TAGCTGGGCA ATAGCTTGAA CCTGGGAGAC CCTGACCTTA TGGGCATCTC TGATATTTGT	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC CATGTCAATC	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA AAAATATGTT AATAGAACTC	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA CATTCTTCAA	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT CACCAAGTCT GCAGCTTGGG	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CAAGGTGGCT  GGGTCAGGGG AAAAAAAAA AAAAAAAAA TAGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC CCAGGAATTG	
10 10 10 10 10 10 10 10 10 10	08601 08661 08721 08781 08841 08901 08961 09021 09081 09141 09201 09261 09321 09381	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACCTGTA TTCGAGACCA TAGCTGGGCA ATAGCTTGAA CCTGGGAGAC CCTGACCTTA TGGGCATCTC TGATATTTGT TGCAATATTG	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC CATGTCAATC TTTGTCCTGA	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA AAAATATGTT AATAGAACTC GCTTCTTACA	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA CATTCTTCAA ACTTTCACCC	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT CACCAAGTCT GCAGCTTGGG AATGCAGTCA	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CNAGGTGGCT  GGGTCAGGGG AAAAAAAAA AT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC CCAGGAATTG GCTCTGTTGA	
10 10 10 10 10 10 10 10 10 10 10 10	08601 08661 08721 08781 08841 08901 08961 09021 09081 09141 09201 09261 09321 09381 09441	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTTGGA CCTGGGAGAC CCTGACCTTA TGGGCATCTC TGATATTTGT TGCAATATTG AAATCAATCA	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC CATGTCAATC TTTGTCCTGA GAATACCTTT	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA AAAATATGTT AATAGAACTC GCTTCTTACA CATTGTTTC	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA CATTCTTCAA ACTTTCACCC TTTGCTGCTT	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT CACCAAGTCT GCAGCTTGGG AATGCAGTCA CTCTAGGAGC	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CNAGGTGGCT  GGGTCAGGGG AAAAAAAAA AT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC CCAGGAATTG GCTCTGTTGA AAGCTGCCAT	
10 10 10 10 10 10 10 10 10 10 10 10	08601 08661 08721 08781 08841 08901 08961 09021 09081 09141 09201 09261 09321 09381	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTTGGA CCTGGGAGAC CCTGACCTTA TGGGCATCTC TGATATTTGT TGCAATATTG AAATCAATCA	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC CATGTCAATC TTTGTCCTGA GAATACCTTT	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA AAAATATGTT AATAGAACTC GCTTCTTACA CATTGTTTC	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA CATTCTTCAA ACTTTCACCC	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT CACCAAGTCT GCAGCTTGGG AATGCAGTCA CTCTAGGAGC	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CNAGGTGGCT  GGGTCAGGGG AAAAAAAAA AT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC CCAGGAATTG GCTCTGTTGA AAGCTGCCAT	
10 10 10 10 10 10 10 10 10 10 10 10	08601 08661 08721 08781 08841 08901 08961 09021 09081 09141 09201 09261 09321 09381 09441	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTTGGA CCTGGGAGAC CCTGACCTTA TGGGCATCTC TGATATTTGT TGCAATATTG AAATCAATCA	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC CATGTCAATC TTTGTCCTGA GAATACCTTT	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA AAAATATGTT AATAGAACTC GCTTCTTACA CATTGTTTC	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA CATTCTTCAA ACTTTCACCC TTTGCTGCTT	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT CACCAAGTCT GCAGCTTGGG AATGCAGTCA CTCTAGGAGC	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CNAGGTGGCT  GGGTCAGGGG AAAAAAAAA AT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC CCAGGAATTG GCTCTGTTGA AAGCTGCCAT	
10 10 10 10 10 10 10 10 10 10 10 10	08601 08661 08721 08781 08841 08901 08961 09021 09081 09141 09201 09261 09321 09381 09441	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTTGGA CCTGGGAGAC CCTGACCTTA TGGGCATCTC TGATATTTGT TGCAATATTG AAATCAATCA	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC CATGTCAATC TTTGTCCTGA GAATACCTTT	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA AAAATATGTT AATAGAACTC GCTTCTTACA CATTGTTTC	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA CATTCTTCAA ACTTTCACCC TTTGCTGCTT	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT CACCAAGTCT GCAGCTTGGG AATGCAGTCA CTCTAGGAGC	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CNAGGTGGCT  GGGTCAGGGG AAAAAAAAA AT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC CCAGGAATTG GCTCTGTTGA AAGCTGCCAT	
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10 10 10 10 10 10 10 10 10 10 10	08601 08661 08721 08781 08841 08901 08961 09021 09081 09201 09261 09321 09321 09381 09441 09501	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTGGGCA ATAGCTTGAA CCTGGGAGAC CCTGACCTTA TGGGCATCTC TGATATTTGT TGCAATATTG AAATCAATCA GGCGGTTTGT CCCTGTCATA	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC CATGTCAATC TTTGTCCTGA GAATACCTTT CTGAATGACC  CAGTTTTTC	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA AAAATATGTT AATAGAACTC GCTTCTTACA CATTGTTTC ACAGTGACCC  TCTATCCAGC	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA CATTCTTCAA ACTTTCACC TTTGCTGCTT CAAACTGGTC ATCAACAGTG	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGAATGTATCT CACCAAGTCT GCAGCTTGGG AATGCAGTCA CTCTAGGAGC TTTGTTTTCA  ATCCTTTTTG	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CNAGGTGGCT  GGGTCAGGGG AAAAAAAAT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC CCAGGAATTG GCTCTGTTGA AAGCTGCCAT CTTTTAATCC	
10 10 10 10 10 10 10 10 10 10 10 10 10 1	08601 08661 08721 08781 08841 08901 08961 09021 09081 09261 09261 09321 09321 09381 09441 09501	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTTGGAA CCTGGGAGAC CCTGACCTTA TGGGCATCTC TGCATATTTGT TGCAATATTG AAATCAATCA GGCGGTTTGT CCCTGTCATA GTCCACTGTC	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC TTTGTCCTGA GAATACCTTT CTGAATGACC  CAGTTTTTC TGCTGAAAAG	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA AAAATATGTT AATAGAACTC GCTTCTTACA CATTGTTTC ACAGTGACCC  TCTATCCAGC ATTCCACTGG	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA CATTCTTCAA ACTTTCACC TTTGCTGCTT CAAACTGGTC  ATCAACAGTG CTTTCCATCA	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT CACCAAGTCT GCAGCTTGGG AATGCAGTCA CTCTAGGAGC TTTGTTTTCA  ATCCTTTTTG CCTTCATAAT	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CNAGGTGGCT  GGGTCAGGGG AAAAAAAAA AT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC CCAGGAATTG GCTCTGTTGA AAGCTGCCAT CTTTTAATCC  AAGGTATTAT AAAAACCAGC	
10 10 10 10 10 10 10 10 10 10 10 10 10 1	08601 08661 08721 08781 08841 08901 08961 09021 09081 09261 09261 09321 09321 09381 09441 09501	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTGGGCA ATAGCTTGAA CCTGGAGACC TGGGCATCTC TGCATATTGT TGCAATATTG AAATCAATCA GGCGGTTTGT CCCTGTCATA GTCCACTGTC ATCCTTATCA	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC CATGTCAATC CTGAATGACC  CAGTTTTTC TGCTGAAAAG TAGCCTACAA	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CAGTGTCTCT CTCATATACA AAAATATGTT AATAGAACTC GCTTCTTACA CATTGTTTC ACAGTGACCC  TCTATCCAGC ATTCCACTGG GTAAGATGAC	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA CATTCTTCAA ACTTTCACCC TTTGCTGCTT CAAACTGGTC  ATCAACAGTG CTTTCCATCA CAACCATTAC	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT CACCAAGTCT GCAGCTTGGG AATGCAGTCA CTCTAGGAGC TTTGTTTTCA  ATCCTTTTTG CCTTCATAAT AGTTTGCCTG	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CNAGGTGGCT  GGGTCAGGGG AAAAAAAAT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC CCAGGAATTG GCTCTGTTGA AAGCTGCCAT CTTTTAATCC  AAGGTATTAT AAAAACCAGC ACTCTCAGGG	
10 10 10 10 10 10 10 10 10 10 10 10 10 1	08601 08661 08721 08781 08781 08841 08901 08961 09021 09081 09141 09201 09261 09321 09321 09381 09441 09501	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTGGGCA ATAGCTTGAA CCTGGGAGAC CCTGACTTT TGCAATATTG TGCAATATTG AAATCAATCA GGCGGTTTGT CCCTGTCATA GTCCACTGTC ATCCTTATCA GTTCCTCAGG	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC TTTGTCCTGA GAATACCTTT CTGAATGACC  CAGTTTTTC TGCTGAAAAG TAGCCTACAA GTGTAAGACT	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA AAAATATGTT AATAGAACTC GCTTCTTACA CATTGTTTC ACAGTGACCC  TCTATCCAGC ATTCCACTGG GTAAGATGAC TACAGTGCTG	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA CATTCTCAA ACTTTCACC TTTGCTGCTT CAAACTGGTC  ATCAACAGTG CTTTCCATCA CAACCATTAC AAACTTAGAA	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT CACCAAGTCT GCAGCTTGGG AATGCAGTCA CTCTAGGAGC TTTGTTTTCA  ATCCTTTTTG CCTTCATAAT AGTTTGCCTG AGTTCCAAGC	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CNAGGTGGCT  GGGTCAGGGG AAAAAAAAT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC CCAGGAATTG GCTCTGTTGA AAGCTGCCAT CTTTTAATCC  AAGGTATTAT AAAAACCAGC ACTCTCAGGG AAACTAGGAT	
10 10 10 10 10 10 10 10 10 10 10 10 10 1	08601 08661 08721 08781 08781 08901 08961 09021 09081 09141 09261 09321 09321 09381 09441 09501	TACTCTGGTG CTCACCACTT AGCTGTAAGT GCTGTTTCTC CCACTTCAAG  CACGCCTGTA TTCGAGACCA TAGCTGGGCA ATAGCTTGAA CCTGGAGACC TGGGCATCTC TGCATATTGT TGCAATATTG AAATCAATCA GGCGGTTTGT CCCTGTCATA GTCCACTGTC ATCCTTATCA	TACTCTGGCT ACATGCTCTG GCCACGTTAA ATTCCCGGGA CTTCAGAACG  ATCCCGGCAC GCCTGGCCAA TGGTTGCGGG CTCGGGAGGC AAGAGTGAAA AATCTCTAGA GAACTTGTCC TTTGTCCTGA GAATACCTTT CTGAATGACC  CAGTTTTTC TGCTGAAAAG TAGCCTACAA GTGTAAGACT ACCTACTAGA	CAGTTCTTGG TGCTCTGTCA CTTCTAGCAA TGCCTAACCA AAACATAGAG  TTTGGAAAGC TATTGTGAAA CGACTGTAAT AGAAGTTGCA CTGTGTCTCT CTCATATACA AAAATATGTT AATAGAACTC GCTTCTTACA CATTGTTTTC ACAGTGACCC  TCTATCCAGC ATTCCACTGG GTAAGATGAC TACAGTGCTG TCTGTACTCT	ACTCCTCTTT AATAGTTTGT TGCCAAAGCC CCTCTCCAAA CTTAAGAAAT  TGAGCCTGGT CCCCGTCTCT CCAAGCTACT GTGAGTTGAG AAATAAGTGT ACTGCATATT TATACGTAAA CATTCTCAA ACTTTCACC TTTGCTGCTT CAAACTGGTC  ATCAACAGTG CTTTCCATCA CAACCATTAC AAACTTAGAA GGCTACCCTC	TCTTGGCTAA TCAACAGAAC TGTGATAGTG TTCTATCAGT ATAGGCCCGG  GGATCACCTG ACTAAAAAA CGGGAGGGTG ATCGCGCTAT TTGCAATTAT TGATGTATCT CACCAAGTCT GCAGCTTGGG AATGCAGTCA CTCTAGGAGC TTTGTTTTCA  ATCCTTTTTG CCTTCATAAT AGTTTGCCTG AGTTCCAAGC TGACCTCATT	CTCAACTGGC ACCACGGCCT GCAGCTTCGG TTGCTTCCAC CNAGGTGGCT  GGGTCAGGGG AAAAAAAAT AGACAGGAGA TACACTTAGG AAACCATCTC AATTGAATAA GTTCTTCCTC CCAGGAATTG GCTCTGTTGA AAGCTGCCAT CTTTTAATCC  AAGGTATTAT AAAAACCAGC ACTCTCAGGG AAACTAGGAT CTCTTCGCAG	

109921 CCTTTATATC TATTCTTTCT CCCTAGAAGG GTCTTGTCCT GGATATCTGA ATGGCTCTAG
109981 ATCTCATTTC ATTCAAGCCT CTCCTCAAAT ACCAACCTTA CGAAAGAGC CTCCCATAAT
110041 CATCCCTTGT AAAATAAGCT TTTCTGCTCA TTTAGCATAT ATATATATAG TTGACTATCC

TCAATAGCAT ATATATAA CATTTCCCCA CCTAGAATTA TATATGTAAT AATATATTA

110161					TGTTCTCTCT	
110221					AAGTATCCCT	
110281					GAATGAAAGG	
110341					AATGCTAACA	
110401					GCGTGCAGTT	
110461					TTATCTTTGT	
110521					GGTGGGTATG	
110581					GAAAGGCCAG	
110641	ACAGGAGCTA	GTCGTGGCTT	ATTGTTACAA	CGACTATACC	TCCCATATGG	GTAATCGATA
110701					GGAATTAAAA	
110761					TCAATAAATA	
110821					AGGCTCATTT	
110881					CATTGTCAGT	
110941					ATGAAAACTG	
111001	AACAGGACCT	CAAAATAAAG	AGACATCCAT	CACTGAAGCT	<b>AACATCGTGA</b>	GGCTGAAATC
111061	AGTCCTATAA	CAATGGTACC	AAAAAGAGCA	CAATGAGAGG	CATTTGTGAA	TATTTACTCA
111121	GATGAGAGTA	AGATATTTCC	CTATCAGCTA	ACCTGAAGTT	CACATCCCTT	TTCCAGCTGA
111181	GTTCTGAAGC	TAGATGTACT	TAACTGGAAC	ACATAACTGC	ATCAGGAACA	TCCTTTAAAA
111241					CAGGTTTAGC	
111301					CTACCACCCT	
111361	TTTCTCTCTC	TGCATATATA	AAAATATATG	TGTATGTATA	TAATCAGCTT	TATTGATATT
111421					TGAATTTTAC	
111481	TAGTTGTACA	ACCATCATCA	CAATTTAATT	TCGGAATATT	TCTATCACCC	AAATTTCCAT
111541	TTCTGCGTAA	AGGGGGAAAA	AAAAAGGTTA	ACTGCTGAAG	GCCGCGGTAA	CACTGAAAAA
111601	GGTGCCTTTT	CTCTCTAAAA	CAGATTTTAA	TCTCCCCTGA	ATTTAGTGTC	CTGGGTATTC
111661					ATAGAGTAAA	
111721					GATACTTAAA	
111781	ATTTTATTTC	ATAATCGCTA	AAAGATGGTT	TTTTTTTTC	CTAAAACAGG	GTTTTTGTTT
111841					CTTGCCTCAG	
111901					ATTGATTGGG	
111961					TTTTCCACTC	
112021					GAGGGCAAAC	
112081					ATCTTAATTA	
112141	TTCTTGACTT	AATGGATACA	GCTCTTCTTT	TGTTTAGTTG	GGCGGCCCTG	AAAAGGGCCT
112201	TTGGTTCAGA	AATGCAAGCT	GTGGAGAAAT	CAGCAACCTT	AACCGCCAAA	GCCATAAAGG
112261	GTGCGTCCCT	GGCGCTTAAG	CGCGTAGACC	ACGTCCATGG	CAGTGACTGT	CTTGCGCTTG
112321					CCAAAAACAC	
112381	CCGCGAGTCT	CCTCGTAGAT	CAGACCAGAG	ATCCGCTTCA	CACCGCCACG	CCGGGCCAGA
112441					GCAACACCTT	
112501					GTCCAGACAT	
112561					TTTATATATC	
112621					CAGGCGGGAA	
112681					ATGGCGGGAG	
112741					CCAATGGCCT	
112801					ATGTAAACCC	
112861					CTCTGTCGCC	
112921					CAGGTTCAAG	
112981					CCGTCGCGCC	
113041					GGCTGATCCC	
113101					AATTACAGGC	
113161					CATTAAAACG	
113221					TACTTTACTT	
113281					CACAGTAATC	
113341					ATAGTGACGC	

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113401	GATGGCGCAC	AGGTTAGTGT	CCTCAAATAG	CCCTACCAAC	TAGGCCTCGC	ACGCCTCCTG
113461	CAGAGCCATC	ACAGCGGAGC	TCTGGAAACG	CAGGTCTGTT	TTARAGTCCT	GCGCAATCTC
113521	GCGCACCAGG	CGCTGGAAAG	GTAGTTTACG	AATAAGCAGT	TCAGTGGACT	TCTGATAACG
113581	GCGGATCTCG	CGCAGAGCCA	CGGTGCCCGG	CCGGTAGCGG	TGGGGCTTTT	TCACGCCGCC
113641	GGTGGCCGGA	GCGCTTTTGC	GGGCTGCCTT	AGTGGCCAAC	TGTTTGCGTG	GCGCCTTGCC
113701	ACCAGTAGAC	TTCCGAGCAG	TTTGCTTAGT	GCGAGCCATC	ACGGAAAAAC	AGCACAGCGG
113761	AACACCCAAC	ACTAGCGCAA	ATACGCCCAT	GAGCTGCTCT	TTTTATACTC	TGTAAAGTGC
113821	AGTGATTGGA	TGATAGAAGA	CGCTAAATAT	GACGTTACAC	, <u>ערהרהראשה</u> הדודאזאפוה	GTCTATCTTT
113881	AAGCCAGCAA	CAATCGTGCA	GTTTCACCGG	CTACTATATA	CTATTCALIG	TCTACAGATG
113941	ATTATTTAAG	TGGTATTTTA	TTACTACTAT	TATTTTATATI	CIALICCAAC	TTGTTCCCCA
114001	AGCTGGTCTT	AAACTTGGGC	TCAAAGGATC	TTCCCGCCTC	TACTITIGET	GTAGCTGGGA
114061	TTACAGGGGA	GCCCACTGC	GCCGGCTTGG	בררכבטבטבעוע	TTTTA A A CTTTC	TCCTCTTCTA
114121	CATCTGGTTT	TCATAACCTG	AAGGCTGTGT	TTATTTTCCA	TARACIIG	CATTGATTCC
114181	AAAGGTATTA	TAATTCCCCA	ATTCCGTATA	ACCUTCACCA	CTTTACCAA	AAAAAAAAA
114241	AAAAAAAAA	GAGGGAATAC	TGCTCACCTC	CTCTCCGGAA	ATCTACCOM	TACGGGAATT
114301	TCTGAAACCT	TTCACAAGAA	TTGGATTCCT	TTGTAATGCT	TTARTCCCII	TAGGAGTGTT
114361	ATTGAAATCT	ACAAAGCATC	TCAAACATAG	TAGGATTAGA	CTATTACT	GAAACATTTT
114421	CTATGAGACG	TCTTTCTCTT	GATTATECTC	TAGGATIACA	CTATTACTCA	CGTTCTGCAG
114481	CTTTTGTTT	CTAAAGCCTA	GGTGTACTCT	CCCACTCACA	AAATGGCGTT	CGTTCTGCAG
114541	TGCCGCCAGG	TACCACCAGC	TGGGAGTTGT	TOCTOTOGO	GAGCAGGAGG	TCTCCAGCAC
114601	CCAAGAGAAA	CTGGATAGTG	GTTCGCAAGG	A A C A TT A STORT	ACCAMMOGAGG	AGAGCTAATG
114661	CAATCATTTT	GAABATCTCA	DADCACTGAA	AACATAATII	AGCATTGCCA	AAATTCACAA
114721	GAGACAGGCC	ACATTCTATC	TTTTGATTGG	TTTNCCCTNT	TTTCTTGAAC	AAATTCACAA
114781	AAAGCAGATC	TATCATCCTT	CATTTCCATC	GACCOTTCCC	ATTTTATTTG	AGCCATTTAG
114841	AACCCAATAG	AAAAAAGGGA	GGCAGAACCC	ATTROUBLE A	GTGGAAACTC	AAACCAGTTT
114901	TAATTAGGAG	TATTTCCTTT	TCARROTTC	CCTTTTTTTAAA	GATACCTCGC	CTGAATCAGA
114961	AAGAAAGGTT	TATATCTTC	ACA A ACCOUNT	TACTORAGE	AATCTTCCAA	TTATTACACT
115021	CTGTGTTTCA	TAACTGACTA	GCCGTCNAAC	CARCARCTAC	AGTTTCCAAC	TTTTGTATAC
115081	CAAATTTTTA	GARATTACGT	GAAATATTTG	AATGCATCCC	TTCTCAATAA	CGTTATTTTC
115141	AGGAAGCACT	GGTGCAGAAG	ATGGGTACAA	TACTTATCTC	GGACCACTCC	AATGGGACGT
115201	TGGCACGTTG	TTTGAACAAA	AAGGGGAAAA	CCTCACCTTA	CTTAGCATGG	ATTATTTGGT
115261	TTTGAAAACT	ACCACAGCAG	GAGCGGAAAT	DAGACCCCAM	TACCTCACTC	TTCGGACTTA
115321	TGTGCTAGGG	GGTTATCCAG	AATAGGATTG	TAGAACCCCAI	TGTCGATTTA	TCTGCTGTGC
115381	ATTCTCCCAT	TAGCTGAGTC	TCTGATTGGC	AATGTGAGAT	CGTTTTAGCT	ATAGTTTTT
115441	TTGAAATGCA	CTTAACAGCC	ACAAACAAGT	TARACCCTTC	TTACCATAAA	TATTGATACT
115501	CAGGGTGTGC	ТТССАТТТАТ	CPCCCCCCCCCC	TOCTOTO	CTAAGTGGAC	ATCTTATCCC
115561	AGCAGAATGC	CTGTCAGGGA	ACCCCATATI	TOCACA	CTAAGTGGAC	TTAACTCCCC
115621	TTGTGAGGCC	CATALATATT	TCTTCAATAA	AACAAMCACM	ATTTAACGCC	TTTCGCAGGC
115681	GATTGCGTGT	GCTGACATGG	AACACAGGTT	CTANACCTOR	AGACCATGTC	ATGGTGCGCT
115741	GTATGGATGA	AAAGGGCATT	GGAAATTCCT	GIAMACCIIA	CCACAMMOCA	GGGGCATGTT
115801	AAGTTGCAAG	TGCAGAAACG	TTTCCACACT	TCCACTTCA	CUACATTEGA	CTGTGGAAAT
115861	GAATTCTGGT	GTTGTCTACG	ATTCATCT	CTTTCACCTC	DARGERAND	CAGCGTTTGT
115921	TCGCTCTAAA	ACATTGCCAG	AAAATGTAAT	DENETTED TO	AMAGGIATIC	GCGAGACACA
115981	CTAAAACTCG	CACTTTTCTC	TOCOTOCO	ACTA TOTAL	ACAACTGGCC	CTAACACGGC
116041	TGCAAATTAA	AAACTAACAT	CTCTGGCAAC	CCACCTCTAA	ACACTGTATT	TTACATTTCT
116101	CGGATGCTTG	TGGCACTGCA	TTTCTABACC	GCCCCCTCTAA	AAATTTCTAA	TAAAACTCCT
116161	GCTGCTTTTT	GAGAGAGAAG	CGGTACCCTC	TGATGTTAGT	CCCCCCACT	CTAAAAAAGA
116221	TTTCCTTCAC	AATGAGGCAA	CCAGAGCGGC	TOWIGITMCT	TOTTTOTTO	CIGCCTACAA
116281	AGCAGGACCA	TAGGCCCTAG	AGGCCCCCAG		AGACTCCCCC	N N N C C C C C C C C C C C C C C C C C
116341	CAGCGCGCAG	GGGGCGCTAG	GGCGCGAGGG	GCGGGCACTC	VCCCCCVCC	ATCACCCCCC
116401	AGTCCCACCC	TATAAATAGG	CTGCGTTGGG	CCCCCVCIG	CCCATCCTCC	WICHCROSCOC
116461	TTATACCACT	TTATTTGGTG	TGCTGTGTTA	GTCACCATGT	CGCMICCIGC	CCCTCCCCC
116521	CCCGCCGCTT	CTGCTGCTCC	TGAGAAACCT	TTAGCTGGCA	DEBRECCARA	CAAACCTCCCCCC
116581	AAGGCTGCAG	CAGCCTCCAA	GAAAAAACCC	COTCCTGGCW	CCCTCTCACACA	COMONHOCHE
				octored I	CCGIGICAGA	GCIGHICGIG

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116641				GGTGTGTCGT		
116701						GGGCATTAAG
116761						GGGTTCCTTC
116821						GGTGGCTACA
116881						TAGCAAAAAG
116941				CCTGCGGCAA		
117001				AAAGTAGCTA		
117061				GTGACGAAGC		
117121				TTAGAAGTTT		
117181				AAGAGCTGTA		
117241	AAGCAAATGC	AACACGCCCT	CAATTATATT	AGAATCACTT	GGAGAGTCGA	TAGAACTTTA
117301	ACATAGCCTC	ATCTAGTAAG	AATTTACTAC	TCAATCTATC	AAAGATAGCA	AGGTGAATTC
117361	AAATGCACCG	AGTTAAAATC	GAGTTTTAAA	GTCACCTGGG	TTTCGGTAGC	CGGAAGTCCC
117421	GCGTCTCACG	ACTCCAAGCT	AATTAGTCAT	AACCGTATTG	AACCAAGGTT	GAAGCCCAGT
117481				TTAAAGTGGG		
117541	ATATTGCTAA	AGTAGCATTT	TCCGAAATTG	GGTGGTCCTA	AGAAATGCTT	CTGGGATAGT
117601	TGGCAAAATA	TATGGCTTAA	CCACGCCCTC	TCCACAGGAG	TGGCTAGCGA	GCTGTCTGTC
117661	CTTGGGAAGG	ACGGTGACCC	TGCTGGCGTG	GCTGGCGCCC	ACGTTGGCGT	CCTCTGAAAG
117721	CCCCGCCAGG	TAGGCCTAGC	TCGCTTGCTT	TCTGCAGCGC	CATCATGACA	AAGCTTTGAA
117781	ACGCAAAATG	CTTTCTTTGT	GCAGCGCCTT	ACCATGGGTG	CACTTACGGG	CTGTCGACTT
117841	GGTTTAGGCC	CTTGTCAGGA	CAAAGGAGCT	TAGTTTGTTG	GAGTTTTAGA	GCTGCAACCC
117901	AAAATCCCTT	GCTCGGTTTC	TCTGTTTTTA	GAAACGGAAG	CGCCCTGATT	GGATATTTGA
117961				ATCAGTCGTG		
118021	GAGCCCACAC	ATTCAAAACT	GAAGATCCTT	TTCTCAGAAC	TGCCCCTTTA	AGCTTTTGCA
118081	ATTTTAATTC	TGGGGGTCAG	ATTTTAATAA	TTGGACTTTT	TTGTTTACAT	CTGACAAGAG
118141				CTTAGTGCAG		
118201				GTTGTATTTT		
118261				GTCAGGGATA		
118321				CTAGGTTTAT		
118381	ACCTATTTTT	GTGAGAAATA	TACATGTTTT	TCTTTGAACT	AAGTATTTTA	CATACACCTA
118441	TCTATATACA	TGCATACTTG	TGGTTTTGTT	TTTTTAAAAA	AAAAAAAAA	AAAACACGTT
118501				CCAGACTGGA		
118561	CACACTGTAA	CCTCCAACTC	CTGGGCTCAG	GCTATCCTGC	AGCCTCAGCA	TCCGGAGTAG
118621				GGCTTTTTGT		
118681				AATGGCCTCA		
118741				CTTGTATGCA		
118801	TAGGGTTATT	AATTTAAAA	ACAAAGCCTG	GACGCAGTGG	CTCACATCTG	TAATCCCAGC
118861	ACTTTAGGAA	GCCAGATGGG	CAGATTACTT	GAGCTCAGGA	GTTCAAGACC	AGCCTGGGCA
118921	ACATGGTGAA	ATCCCATCTT	GACAAAAAAT	ACAAAAAATT	AGCAAGGCCC	AGTGGCACGC
118981	ACTTATAGTC	CCAGCTACTT	GGGAGGCTGG	GGTGGGAAGA	TGACTGGAAC	CTGGGAGGTA
119041	GAGGCTGCAG	TGAGCAGAGA	TCGTGCCACT	GCACTCAAGC	CTAGGTGACA	GAATGAGACC
119101				TTACAACGAT		
119161				ACCCTGTCAT		
119221				TATTGATAAG		
119281				TCCAGGGCTG		
119341	GTATTTTATT	ATTTTATTTA	TTTATTTATT	TATTTAGACA	GAGTCTTACT	CTGTCACCCA
119401	TGGTGGAGTA	CAGTGCCATG	ACCTCAGTTG	CAACCTTTGC	CTCCTGAGTT	CAAGCGATTC
119461	TCATGCCTCT	GGTCTCCCGA	GTAGCTGGGA	TTACAGGCAC	CTGCCACCAG	GCCTGGCTAA
119521	TTTTTGTATT	TTTAGGAGAG	ATGGGGTTTC	ACCATGTTGG	CCAGACTGGT	CTCAAACTCC
119581				CCCGAAGTGC		
119641	ATGGCGCCAG	ACCTGGACTT	TGTCTTCTGT	TTCATCAGTC	CTTCTGTTGG	TTCAAGCACA
119701				AAATATGGTA		
119761	TTCTTATTTT	TTAATTTTAA	GGCAATTTTA	GATTCCAGCT	TTCCAAAGAA	TTGTGGAATG
119821	CTTAGAGCTA	GAGAAGCCTT	GGAAGTCATT	TAGTTTTTGT	TTTGTCAGAG	AAAATTCTGT

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	119881				ATCCCATAGT		
	119941	GGCAATAATA	CCTTATGGAC	AGTATGCTTT	TCCTCAAATA	TATTCTAAGC	CATGGTCAAT
	120001	GCAAAAGAGT	GAGAAGGAAA	GTAGAATAAG	TTATCTAAGA	ATCAGTGGGT	GCTCTCTTTA
•	120061				TCTTGAAGGT		
	120121				TAAGTTATTC		
	120181				CTGCTACCAT		
	120241				TTGAGAGGGA		
	120301				GATGTTATGC		
	120361	AGGAGACATG	TTGAAAGTGA	CCCATAAATC	TGCAGTATCT	CATGTCTCTC	AATGGGGACA
	120421	AGGAGTACCA	TGGGAAATAG	CATTAGGTCA	ATGACAGTAA	CAACTCCCAG	GTGAGTTGAT
	120481				GCTACATAGT		
	120541				GATAAATGAA		
	120601				ATTGGAGATT		
	120661	TTAAAATCCT	TCTCATTGAA	AAAAAATTTC	GTATTAGAAG	ATTTCAACAT	TCTTTAAACT
	120721				AGCAAAACAA		
	120781				ACACAAGTAA		
	120841	CCCTCCCAGA	GCCGCCTGAT	GCTTGCTTCC	AGTCACATTA	TCACTCCATC	TGCCCTAAAC
	120901	ATAACCCCTA	TTTTGATTTC	CAATGCTGTA	ATTTAGTATG	CCTGTTTTTG	AAACATATAA
	120961	AATGGAAATA	AAACAAATGT	AATCCTATGT	ACCTGACATA	TTTCACTCCA	GAACATTAGG
	121021	TTTGAATAGA	TTCATCTGTG	TTGCTGTGTA	TAACTTTAAT	TCATTTTTAT	TGTTATGTAA
	121081				TGTCTACTGT		
	121141				ATTCCTGTGT		
	121201				AGGAATTGTT		
	121261				GATGAAATGA		
	121321	AACAGCATGG	ATGAACCTCA	CAAACCTAAT	GTTGATGGAA	TCTAGCTGGG	AATTCCTGTT
	121381				ATTAAAATTG		
	121441				ATTATGGTTT		
	121501				ACTTGGGCTG		
	121561				CTCACAGTTC		
	121621				TAGTGAGGGC		
	121681				CAGAATACCA		
	121741				GGCTATAAAG		
	121801				AGATGTGAGA		
	121861				CCTCTTCATA		
	121921				TCTATCAGAC		
	121981				CCCAAAATCC		
	122041				GGGGGAACAC .		
	122101				CTTTTTGTCC		
	122161	TTGAGGTGGA	CTCTTGCTGT	GTCACCCGGG	TTGGAGTGCA	GTGGCGCTGT	CTCAGCTCAC
	122221	TGCAACCTCC	ACCTCCTGGG	TTCAAGAAAT	TCTCCTCCTC	CCAAGTAGCT	GGGACTACAG
	122281	GTGCATACCA	CCGCGCCCTG	CTAATTTTTG	TATTTTTGAT	AGAGACGGGG	TTTCACCATG
	122341				CAGGTGATCC		
	122401	ATGCTGAGAT	TACAGGTGTG	AGCCACCAAA	CCTGGCCTGT	CTTTTCTGTT	TTAAGTTTTT
	122461	AAATTTTGCT	CACGAACCCT	TTATCCATTT	TATGTGTTGC	AGGTATTTCC	TCTGTAACTT
	122521	GTCTTCACTC	TGTCAGAGGC	TGGAGTGCAG	TGGCACAATC	ACAGCTCACT	GCAGCCTCCA
	122581	CCTCCCAGGA	TCAAGCGATC	CTCCCATCTT	ATCCTCCTTA	GTAGGTGGGA	CTACATGTGC
	122641	AGGCCACCAT	GCCCAGCTAA	TCTTTGTATT	TTTTTGTAGA	GATGGTGCTG	TTGCCCAAGT
	122701	TGGTCTCAAA	CTCCTGAGCT	CAAGCAATCC	ATCAACCTTG	GCCTCCCAAA	GTGTTGGGAC
	122761	TAGAGGTGTG	AGCCACCACT	GCACCCAGCC	AATGATATCT	CATGATGCAT	TAAAGTCATT
	122821	AATTTAGTGT	ACTCAAATTA	AGCACACTGC	CCTTTTATGC	ACAACCTTTT	TTGTATCTTA
	122881	TAAAAAATTT	CATTTTCTAT	TTCAAGGTCA	TGAAGATCTT	ATTTTATAAT	ACCTTCTTGT
	122941	GAAATTAGTT	CTCAAGACTA	CCCTCACTTC	TAACACCAAT	TATAAGTTGG	GAGGTCTGTG
	123001	GTTCCCAATC	AACCTTAGGT	TAGTAATTTG	CTAAAAGGAC	TCACAGAACT	TGCTGAAGCT
	123061	GTTAGCCTCA	TGGTTACAAT	TTATTATAGG	ATATATAGCT	TATTATGTCA	TTCCAATGCA

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123121	ATGTAAAATT	ATACAACTAC	${\tt TTTTAAAAAG}$	ATTTTAGCAT	TTGACCCAAC	AATTTCACTC
123181	TGAGGTATAC	AAACAGCAGA	TATGTGTGCA	CATATATACC	AAGACACATA	CACAGCAAAA
123241	TTCATTGTTT	GTAATAGTTG	AAAAGGGGAA	ACAACTCAAG	GAATAAAGAT	TAAAATCAGC
123301	TGAGAAAAGA	AACACACAAG	GCAGTATTAT	GGATCGAATT	GTATGCAGAT	CTCCCTTGCC
123361	CCCAGAAGAT	ATGTTTAAAG	TCCCAACTCC	CAGTACCTCA	GAATTGTGGC	CTTATTTGGA
123421	AATAGGATAG	TTGCAGATAT	AATTAGTTAA	GATGAGGTTA	TAGTACAGTA	TGATGGGCTG
123481	GTGACTTAGA	AGAAGTAGTA	TATATATATT	TTTTAATAGA	ACTAGTATTC	TTCTAAGGTG
123541	GTCACGTGAA	GACAGACACA	CACAGGCAGA	GACTGAGGTT	ATGCAGCTGC	AGGTCAAGGA
123601	ATGTCAAAGG	TTGCCAGCAA	GTACGAGAAG	CTAGGAAGAG	TCAAGGAAGG	ATTTTCCTAC
123661	AGGCTTCAGT	GGAAGCATAG	ATCTAATGAT	ACCTTCATGT	CAGATTTCTA	GCTTCCAGAA
123721	CTACAAGAGA	ATATATTTGT	TGTTTTAAGC	CACCCTAGCT	TCTAGCTCTT	TGTTACAGCA
123781	GCCCTAGGAA	ACTAATATAG	GCACAATCCA	GGCAAGTTCC	AAATATGAGC	TTCCAGTTGT
123841	CCTCTCCCAG	TAATATGAAC	AGTATTACTT	TCCCAGCATT	AATGTGTGAC	AATACACATG
123901	ACGTACAGAG	CAGTCCCCAC	TTATGCACAA	AACATATGTT	CCAGGACCTC	CAGTGGATGT
123961	CTGAAACCAT	GGATAGTACT	GAACTCTATA	TAGCTGTTTT	TTCCTATACA	GACACAGCTA
124021	TGATAAGGCT	TAATTTATAA	ATTAGGCACA	GTAAGAGATT	AATAACAATA	AATTAGAATA
124081					ATTTCTGAAA	
124141					TGTAGAAACC	
124201					GCTGCCATG	
124261					GATACTGTGT	
124321					ACCTCCAGGT	
124381					CTCCCAGTAG	
124441					ATACTTTATT	
124501					ATTGTTTTTT	
124561					CTTTTTCTGT	
124621					ACCATTGTTG	
124681					TCCATTCGTT	
124741					GATATTTAAT	
124801					GAACCCTGTT	
124861					GTTTTGTAGA	
124921					AATATACTCC	
124981					TTCTTCGTAG	
125041					TAATACCAGC	
125101					CAGACTGGCC	
125161					GGTGGCGGGC	
125221					CCAGGAGGCA	
125281					GAGCGAGACT	
125341					TTCAAATAAC	
125401					ATAACCTCCA	
125461					AAAAGGCTGG	
125521					TCTGGCTTCA	
125581					TCAACAAGGG	
125641					TAAGGGACAC	
125701					CTCTGGAACT	
125761					TGGAGGTTGC	
125821					GAATGTGAGA	
125881					TTTTCTTATC	
125941					AACAAGCTTC	
125941					CTATGTTGTC	
126061					GTTGGGATCA	
126121					GTTATTACAT	
126121					GTGAAGTGGA	
126241					AGATGATGTA	
126301					GAGAGAAAAT	
750307	PHIDMONDAG	CUCTOWWIT	TANAMANAT	-VOVIGCRITI	CHANNED TO	TAGGRAGIAG

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SUBSTITUTE SHEET (RULE 26)

126361	AATCAACAGA	TTAGATGTAG	GGATGAGAAG	GGTCAAAGAT	GACACTAGGG	TTTTTAACTG
126421	GAGCAAGTAG	GTAGACAGAA	CATTTCTTCC	TGAAAGGGCA	GGTCAGATCA	TGTGTTGTCT
126481	CAAAGGGCAT	GAAGAGTAGA	AAGCCTGGGA	CAGATCCTGA	GATGACCAAT	ACCCATGGTG
126541	CAGGGAGAGG	GAGGGAGATC	TGCTAAAAAG	ACTGCAAATG	TCAGGATAGT	AGAAAATCAT
126601	GAGTGTGTGA	TGTCCTGGAA	GTTGAGACAG	TATCACATTT	GAGAACATTT	AAATTGGTAA
126661	CTCTGACAAA	AAGCTGGAGG	CCAACTGTGA	ATGCCCATGA	GAGTGAGAAG	CTCCCACACT
126721	TTTGTGGGCA	TCAGAAAGCC	CACCAGGTTC	CTGCAGTGAA	GATCTGAGAA	GGATCCTCTT
126781	GTGGCTTTGG	CAGGGAGAGA	AGAATTATTA	TGAAATACAC	CCCAGAACCT	TCTTCAAAAC
126841	AAAGGCCTAC	TCTCAAGGGG	AAAACATTTT	GCCAGAGTCT	TATCCCAGCT	GGGAGAAGGT
126901	AATTCTTCCC	ACTGCAGCCT	CATCTAGGCT	TTCTGTCTCA	CTTAAGGGAA	GAAAATTAGT
126961	CAACAGGGAT	CAGAGCTTCA	TGAAAATAAA	TTGGAAATGG	TGCAGCCAGG	AAAGGAGCAA
127021	AGGTCTGAGG	AGGAGGAGAA	GGAGGAAGAG	GAGTTGTATC	ATTATAAATA	CTTGAGGAAG
127081	AGGAGGAGAA	GGAGGAGGAG	GAGGAGTTGT	ATCATTATAA	ACACTTGAGG	AAGAGGAGGA
127141	GGAGAAGGAG	GAGGAGGAGT	TGTATCATTA	TAAACACTTG	AGGAAGAGGA	GGAGGAGAAG
127201	GAGGAGGAGG	AGGAGTTGTA	TCATTATAAA	CACTTGTGAC	GGTCCCAGCC	CCAAGATATA
127261				TGACTACAGA		
127321	CATCAAGGCT	CCAACTGAAT	AACAATGAAT	TATGAATGAA	AGAGCTGTAA	GGAGAGACAA
127381	AAGTTAGAAT	GAGACAAGTA	TTGTTATCTA	GAGATGCCAA	GAAGGCAAGG	AAGATAACTA
127441	AAAAGGCACT	CTGGATTTAG	AAATAGGAAG	TCATTAGTGA	CCTTGTAAAT	AATGGAGCCA
127501	GAGGAATACC	AAGGGCAGAA	GCCTCACTAT	AGTGTGTTGC	ACCTGTCAGA	GGTCAGGAGG
127561	TGTAACTGAC	TCTCCCACAG	TGTGGCTTTG	GAAGAGAGAA	GTCAGCAGCT	GCATGGAGAT
127621	TTGGGAGAGG	GAAAGCTTTT	TTTTTTTTTT	TTTAATTGGA	AAAGACTGAG	CTATGTGTAA
127681	ATAGAATAAG	ACAGGAAGAG	TGTAGACACA	GGAAAGAGGG	CAGACAAAAA	CAAGTGCACA
127741				GCAAGTATAT		
127801				CCAAACCCAC		
127861	TGACTTGCTC	TGTGCCCCAG	AAGCCCAGCT	TCTACAGATA	GCATTAGCTG	GGCAGCCCTG
127921	CCCTCTTGCA	ACAGCTGGAT	TTGGCCAGTG	ATCAGCCCAG	CAGGAATGTA	GATGGCAAAG
127981				ATCACCCCC		
128041	CTCCACAGTC	CCAGCTCTGG	CCTAGCTCTG	GTTACAGGTT	CCCTCCCATT	GCCTCTTCAG
128101	ATTTAAAGGT	GTGTCTGTCA	GGGTATAACT	GGGAGCTAGA	AATTGCACTG	AAATTGAACA
128161	AAGAATTTTA	TGGGAATGGT	TGTTAACTAG	TTATAAGAGG	ACTGAAAATG	GAAAAGTGGA
128221	CAAACGTATC	AGAGATAGTA	ATGACAGAAA	GCAACTACCA	CCTCCAGGTT	TAGGAGAACA
128281				ACTGGGACCT		
128341				CATGATGAGG		
128401	GATCTCCAAA	CTGAAGCCAA	CTGCTGTTAC	TGGATTCAAC	TGCCACTGCC	AGGTTGAAGA
128461	ACCCATTCTG	TGAGGATGTC	AACAAACAAA	GTGGGAAATC	TTTTCACATC	CTTCCAGCCC
128521	TCTAGTCTTC	CTCCAGTGCT	TTCTATTGGT	AGGGTTTGGG	GAGGTGGCTA	GCAAAGCGGT
128581				ATAACCAGCA		
128641	ACTGTTGCTG	ATCTTGGGCT	GCCTCATATC	CCCTGTTCTT	CCCATTAGCC	CTGTCACAAC
128701	TTTGTAGATA	TCCCTTCATT	ATATGCCCTT	CATATATTCT	TTTGGTTTAA	СТОТСАСЛАС
128761	TGGAATCCTA	ATATGGCACT	CCTCCATTTT	TCAGGACCAA	AAGAGTATAA	DAGATTATCT
128821	TTTACCAAAA	AAAAGACAAA	AAACTGATCT	AATTCCTGAT	TTGATCATTA	CACABTCTAT
128881	ACATGTATCA	AAATATCACA	TAGTACCCCA	TAAATATATA	CAACTGTGTC	CATTANANT
128941	AAAAATTAAA	GAAAAGATGG	TAAATATAGC	TCTGTCAGGC	AGTGCAGGTT	TTACCACCAT
129001	GGCTGTTATT	TCCCCCATGA	AGGGGGGAGT	GAGGGAGCAG	CTGARAGTAG	GTGCTTATAG
129061	GGGTATAGAG	GGGCTCAAAG	CTTTGAGAGA	GGAGAATGTC	TCDARGOTA	GCCANATAGC
129121	ATGCAGGTCC	CATGGGGGCA	GAGCCTCTGC	TCATTCACCA	GTGCCTCTTC	AATATCTACA
129181	CTTAAGCCTA	ACACAAAGTG	TGTGCTTAAT	AAGTATTTGC	TCACTATCTA	DACTCCARAC
129241	AGAACCAATC	TGGCAAACTT	TGTAGGACTG	GTGGGCAATG	PECHANICAL	AGGTANN NATC
129301	TGTGGATATA	AATTTATATT	GATCAAAAA	TTCAAGGTTA	CCACACACACACACACACACACACACACACACACACAC	TANNATE OF THE PROPERTY OF THE
129361	CTCAACGATG	CTTCAGCCAT	GCTCAACTCT	TCTGTAGCCA	CDCDDDDDDD	TICAGICAIG
129421	ATCGAGCTGT	GTCTGTGTCT	GAATAATGAA	AAGACCATGA	TGCAAGGGAG	TTCGAGACAC
129481	AGAAACAGTG	TTTGAAGTAA	TGGGTAATGG	AAGCATGCTA	CCACCCAAAC	CDADCADCAC
129541	GCAATAGGAA	GGAACAGAGA	TCTGTGGTCC	TATGTCCCCT	CACCATATTC	ACATCTTAAA

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129601	GCTAATTCAG	TTTTCAATCA	TCATTAAAAT	TTTGTTCCTA	AATATATGGC	CATTATTTTC
129661	CACAACCACA	CTAAAACTTT	ATTACCTCTG	GCAAGTGACT	ATGCAAGTAA	CTAAGAGCAA
129721	AAATATCCAC	AACTACCATT	TGAGCTATCA	ATTTAGGGAA	AGTCATCTGG	CTATAATCTA
129781	AGTGACCCTC	CACTGAATGT	CAGTATCTTT	GCATATGTGA	TTTAAATCTG	GGCCTTCGCA
129841	ACACCATGAA	CTGTTCTTGT	CTTGAATATC	CAGATTGAAG	GAAATAATCT	GAGTAGTTAC
129901	GAGTCCTGAA	GCTAGAAAGA	TGGAAACCCC	ATTTGCTCAT	CAGAAAGCCT	TAGAGCTTGG
129961	GCGCTGGCGG	GTCCTGTCTC	ACCGGGACAG	AGGGGCTCTT	TCCTCCCCAT	CTGATAGTCT
130021	GATAACTAGA	GAAGCCGGCC	AACTTATTCT	CCAAGAAGGA	GCCATCTTAG	TTCCTCCTGA
130081	AATGTTCATA	TTTAGAAATT	ATTGTTTGTC	AGTAATTTAA	CCCCTTAATG	GGCTTGCCTT
130141	GTGGTCCATA	CCACTGAGTG	CAGAGCTTGC	CTGGAAGAAT	TGTGAGGGCC	ATTCCATCTT
130201	CCAGGCAGTA	GAGTTCAGTA	CTTCTTTAAA	ATTGCTGCTG	AACTCTGTAT	TTGAAAAGAA
130261	AGAATCATTT	GGGTGTGGTA	GCTCACACCT	GTAATCCTAG	CGCTTTGGGA	GGCTGAGGTG
130321	GGAGGATCAT	TTGATGCCAG	GAGGACCACT	TGAGACCACC	CTGGGTAACA	TAGCAAGACC
130381	CTGTCTTTAG	ААААААААА	TACAATAAAA	TAAATACAAT	AAAATAAAA	GCAAAAAGAA
130441	AGAGTCCATC	TTAGGGACAG	ACTGTAACTA	CTCACTGGAG	CTTACCTTTA	CATAGTTCAG
130501	GATCAATTAT	AATAAAACAC	TTTTGTGCAG	ATTCAATAGG	ATTATTTTAA	TCCCCATCAT
130561	CTCTCTGAGT	TTCCAGTCAG	TTTCTCTGCA	TGTAGACACC	CTTCTCCAGC	CCACCATTGT
130621	CTCTCCTCCT	ATAGCTCCAC	CAACAAATCA	GAACTTTTTC	TAACTGCACC	TAGTGCACCT
130681	AGAGTCTACT	CCAGAATGCT	CATGGAGAAA	GTTTCTGAAA	GGTAAAACTC	TGAATGATAT
130741	TTGTAGCTAA	AGGGAGACTT	GCTAGAGACA	ATAAGCTAAT	AGTTGTAGAC	TTCAGTAGAA
130801	GAGGAATGAC	ACTGCAATGT	CAGGGTGCAG	GACTTCAAGA	GGGCAGAGTA	TGGAAACCCA
130861	ATGGGAAAAA	TGCTCACCAG	GAACATGAAG	AGAAGGAATT	ACGTGTAAGG	ATTTCTCAAT
130921	GTGTTCCCAA	ATTTGCCCAG	CAGAGGGAGG	CCTCGGGTTG	ATGGCAGGCT	GACCACACAA
130981	TTAAAGAAGG	CTGAACCTGG	GGGCTTTTAA	CAACCATCGT	GGGCTCTACT	GTAAGCATTT
131041	AGAAAAAGAA	AGTTATCCAT	TCAAAAATAT	ATATATTTTT	AAACTTCAGA	ACAAAATTAT
131101					ATCTGAGTAT	
131161					TCACTTAACC	
131221	GATTACTCTG	TGTTCATAAT	AATCACTTTT	TTAAAACTTT	TATTTTTATT	TATTTATTTT
131281					TGCAGTGGCG	
131341	TTACTGCAAC	TTCCACCTCC	TGGATTCAAG	CAGTTCTCCT	GCCTTAGCCT	CCTGAGCAGC
131401	TGGGATTACA	GGTGTGCACC	ACCAAGCCCG	GCTAATTTTT	GTATTTTTAG	TAAAGACGGG
131461	GTTTCACCAT	GTTGGTCAGG	CTGGTCTCCA	ACTCCTGACC	TCATGATCTG	CCCACCTTGG
131521	CCTCCCAAAG	TGCTGGGATA	ATCACTTTTT	ATGCTGCATA	ATTCTTCAGA	TTTGTCAGTA
131581	CGACTGTATT	TACACTCATT	TGTTTTATTA	GAAAGAATTC	CAGAATATTT	TGGCTGCCCT
131641	AATTAATTTT	ACAATTAATA	TGATTTTGAA	ATTGGGTATT	GGCTCCTTCT	GAATTGGTTT
131701					TATTAGCATA	
131761					ATAGCTTGTA	
131821					TCAAGATTGC	
131881					CCTCTTTTTT	
131941	GCATGTTTCC	ATTTTAACAG	AACCCGTATT	TAATCAGCAG	ATTTCTATGG	TGGCTAGATT
132001	TGTAGACTAA	ATATTAAAAG	TCCCAAAGCA	AATGCATTTT	TCTCTTAAAT	TTTACTGACT
132061	TTTTTTTTT	TTCTTTTTCT	GAGACGGAGT	CTTGCTCTGT	CGCCCAGGCT	GGAATGCAGT
132121					CACGCCATTC	
132181					CCCAGCTAAT	
132241					CTCGATCTCC	
132301					CATGAGCCAC	
132361					CATAACGTAT	
132421					GAAGGTTTAT	
132481		,			TGAACTACTG	
132541					CTTATGTTTC	
132601					ACTTTCCTCC	
132661					CTTTGGTTTC	
132721						ATGGGGAAAA
132781			-			AAACCCAGCA

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132841	GGAATTCCAG	AAGAAAACTC	CTCAGGACGG	GCACATTGGC	TCATGCCTGT	AATCCCAGTA
132901	CTTTGGGAGG	CCGAGGTGGG	CAGATCACTT	GAGTCCAGGA	GTTTGAGACC	AGCCTGGTCA
132961	ACATGGCGAA	ACCTCATCTC	TACAAAAAAT	AAAAAAATTT	GTCAGGCGTG	GTGGCATGCA
133021	CCTGTAGTCC	CAGCTACTCA	AGAGACTTAA	GTGGGAGAAT	CACTCGAGCC	TTGGAGGTGG
133081	AGGTTGGTGA	GCCGAGATCA	CGCCACTGCA	TTCCAGCCTG	GGCGACAAAG	TGAGACGCCA
133141	TCTCAATCAA	TCAGTCTCCT	CGAAAAGCAA	CATTATGGAG	AGACAGGATT	CCGTCAAGGC
133201	CTGGGGCACA	CAGGAAAATA	TTAAGGCAGA	AGAGAGTTTC	CTCCCCACAC	CACACCGTAT
133261	CCCACAGGCA	CTGCGGATGT	GCATATGCAA	GAGGGGTTGA	TCCTAAGAAT	TTAGAGTCAC
133321	AGAGGAGGAG	GCACCAAGCA	GACTGTGGAG	<b>AAAGTCATGA</b>	CCAGAAAGGG	ACAGAATGTA
133381	AAGCTTCAGC	TGATTATCTG	GCCTCAGGGA	TTCCAGAGGA	ACTGGTCCCA	ATGGTCTCCT
133441	GGTGATGTAG	GTTCTTAGGT	TTCTTTTACA	GGGGTTTTCT	GGGAGATCGT	TGACCCAGTT
133501	AGCATTCAAG	CAACTTCCAC	CCTGCACTTT	TATTCTTTCC	CCTTCACCTG	CTTAGGTTTT
133561	ATCTGTCCAG	GAAATAATAA	TAAAATTATT	GAGCCCTGGA	CATGTACCTG	TAAAGCTCCT
133621	TAAAGATGAT	GCCTTCTAAC	TCCTCATTCA	ACAGATACAA	AAACATTACA	ATAAAATGAC
133681	TCATGCAAGA	CACCCAGGTA	GTTTATAGCA	GCTAATAAAA	ACAGAATAAC	TATAAAATAT
133741	GGTAAGTTTA	TAAAAGTTAC	ATTGAGTATA	CTTTATAAGA	ACTGCTTATT	GAGTTTGCCT
133801	AATAACCACA	CAGCACAATA	ATAATATGTA	TATATTTTTA	AATATGTGTA	AATATGTGTA
133861	ACACAAACTT	GTAGAAGGTA	TATCTGAGTA	CAACCCTATT	CTGTTTGGTT	ACCTTTTCTA
133921	GTTCATTATG	TAAGTGGCAT	AGCTACCTAA	GGACTTATGC	TTATAAATGT	TACTCAAAAA
133981	AATACAGAGG	ACATATGTGG	ATAGATAATG	GAAGAGATAA	GATAGGTAGG	TTGAAGGGTT
134041				TCGTTAGGTG		
134101				GAAAAAAAGG		
134161				TTCCCTTAGT		
134221				CAAAGGATAA		
134281				ATAAACAAGG		
134341	TTTAGATATG	CATACACATA	AACATCTCAA	TGACTTGAAG	AGCAGTATTG	CTGCCAGCAT
134401	GTCCCACCTC	CAGCCCTAAG	GCAGTTTTCC	CCTATCTCAG	TAGATGGAAT	ATACAATCGG
134461				GACGAGCAGG		
134521				TTTACTAATC		
134581				TTTCCCTTCC		
134641				CTGGCTTTCC		
134701				TGAGATTAGG		
134761				AAAGCACATC		
134821				AGGGAGCACA		
134881				CTTAGTACAG		
134941				AATGTGATCT		
135001				CAAAACAAAA		
135061				GTAGAAAGGG		
135121				TAAGGCAATA		
135181				TTAAAAACAT		
135241				GCCGAGGCGG		
135301				AAAATCCCGT		
135361				GTACCTGTGG		
135421	AGGTGGGAGG	ATCCCTTGAG	CCCCGGGAGG	TTAAGGCTGC	ACTCACCCAT	CATARTCCCA
135481	CTGCATCTCA	GCCTGGGCAA	CACACCCACA	ACCTGTCTCA	ANTONOCCAI	DARAGECA
135541				TACCAGTACC		
135601				GGTTTAAGCA		
135661	GCTAGTGACT	CATTCACTCA	TTABACAAAT	ACTTATTGTG	Chuchychyn	AMGGICTIAG
135721	CTGTGCTAGG	TACABARGON	ב בתתתתתת ב	GCTCTATAAA	CATCIACIAL	THE THACTA
135781				ATTCTGAGCT		
135841	TGGCAGAGCT	TTTTAGGCA	TOTACICATE	AGCTCTACCA	CTCCCTCCTT	CAMCOMCCO
135901	TGCCCAAAGG	ATCCATCCTC	TICICIMMMI	CTGCTTACCT	TACCUCIGGIG	COMMUNICA
135961	ACCGCTCTGC	TCTTCTGCAG	TIGGCCCIGI	PALCALIMOCT.	THOUSE CAR	TOTTLIGUAGE
136021	CTCTGCTCT	TTTGATCTG	THE TOUCH	ATGTATGTAA	ACCMCCMMMC	TCIIWGGCIG
			TATACMICAC	PAIDIAIDIAA	AGGICCTITC	CLIMITIACC

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136081	CATGACCAAG	GTATTATGAG	ATTCTGGAAT	TTCCCCAAAC	CACATTGATT	GCTGGGAGAA
136141	TAGAAGAAGT	GGATTACAAG	TGGAACTTAG	AAGGGGAGTA	TTCGAGAAGA	CGTCTCTGCA
136201	AATCCATTTA	GAGAGACCTT	TCTCCAGTGG	TGACTCAAAG	ATGCAGCTCC	TTTCATCCTG
136261	TGGCTTGGCC	ATCTTCAGCA	CATGGCTCCC	AAGGATGTCC	TCAGGATGGT	CTCTAATCCA
136321	AGGAGCCTGA	AGAGAAAAA	AGGCATGGAG	TATTGTGAGT	GGTAGGTGGT	TATGGACCAG
136381	TTATGGAAGA	ATACACATCA	CTTTTGCCCA	CCTTCTACTA	ACCAGAACTC	ACACAGCCAT
136441	AGACACTGAC	AAGTAGGACT	TAACAAGAAT	CTAATTTTGA	GTCTAGGAAT	ACGACTGTAG
136501	CAAATATTTA	ACAGCTTCAA	ACACAGGTGC	ATTGCTATCA	CTATGCTTGG	CCCAGGCCTG
136561	TCTCCCTTTC	CTGCCATGTC	ACAGGGGCCA	GCATTTATGT	CTAGATTGGG	TTGGTTGGGA
136621	TATTAAGACA	ATAATGAACC	AATACAACAT	CTTGAGCATA	AAACCAACTG	ATACAATGAT
136681	GTACAAGTCA	GATGATTCTG	ATGATTATGA	ATTATGTCAA	TAAAAGAAAT	GTGATAACTA
136741	AGGTAATTTT	TGTTTTGGCA	AATTTTTGTT	TGTTCATGAC	AGGATGAAAT	CCTGTCATTT
136801	GTAGCAACAT	GGATGGAATT	GCAGGATACT	ACATTAAGTG	AAATAAGCCA	GAAACAGAAA
136861	GTTAAACACC	ACATGTTCTC	ACTTATATGC	AGAAGCTAGC	TAACTAAGTA	AATAAGTTTA
136921	TCTCATTGAA	GTAAAAAGTA	CAACAGAGAT	TACTAGAGGC	TGGGAATGGT	AGGGGAAAGA
136981	GATGATAAAG	AGAGATTCGT	TAAAATAAGT	TACAGCTAGA	TAAGAGCAAT	CAGTTCTAGT
137041	GTTCTATTTG	TACTACAGAA	TGGCAATAGT	TAACAGTAAT	AAATAATTTC	AAAGAGCTAC
137101	AAAAGAGGAC	ATTGAATGTT	TCCAACACAA	AGAAATGAGA	AATGCTTGAA	ATTATECTES
137161	TTCTAATTAA	TTACCCTGAT	CTGATCACTA	TACACAGTAT	GTATAAAAAT	ATANIGGAIA
137221	GCTGGGCGCA	GTGGCTCACA	CCTGTAATCC	CAGCACTTTG	GGAGGCCAAG	GTARCACIAIGG
137281	CACTTGAGGT	CAGGAGTTAG	AGACCAGTCT	GGCCAACATA	GTGAAACTCC	ATCCCTACTA
137341	AAAATACAAA	AATCAGCCAG	GCGTGGTGGC	ATGTGCCTGT	AATCCCAGCT	ACTORIZED
137401	CTGAGGCAAG	AGAATTGCTT	GAACCCAGGA	GGCGGAGGTT	GCAGTGAGCC	CARACCCCC
137461	CACTGCACTC	CAGCCTGGGT	AACAGAGCAA	GGCTCTGTTT	CAAAAATAAA	TARATICOCOC
137521	AATAAATATT	TTTTAAAAAA	AGAACATCAC	TATGCACCCC	ATATATACAT	ATARTACATA
137581	TGTCAATTTG	AAACATAATT	TTGAAAAATG	AAAAAATGAA	ACACAAATAT	CARTIALIA
137641	TCTCCAAGTT	GATATACTTA	AAAGGAAAA	AGTCCGAGGG	CTTAAACTAT	TCAATCAATCC
137701	TTTTATTAAA	ATGCTATAGT	AATCTGGAAA	GTATTTCAGA	ATGAATTGGT	TORRICARMA
137761	ACACAAAGAT	CAGTGAAACA	AAACAGAGAA	CCCAGDAATA	GATTCACACA	TCTNTCCACA
137821	ACTGGTTTTG	ACAAAGGTGT	CAAGGCTATT	TAATAAGTAA	AAAAATCGTC	TTTTCACTA
137881	ATGTTTCTTG	AACAAGTAGA	CATCCGGTGT	GGGGGAGAGG	AGCAGGAGCC	TTTTCAGIAN
137941	CTTTATGCAA	AAATTAACTC	AAAATAGACC	ATAGACTTAA	ATGTAAAAGC	TARACTORAG
138001	AAACTTCTTT	AAAAAATAGG	AGAAAATCAT	CAACACCCTA	GGATTAGCAA	TANAMITAL
138061	AAAACAAAAC	AACAGGTTTA	TAGTTTATAA	BACATAAATA	ACAAAATGAT	AGRITICITI
138121	AAAAGTGAAA	ATTTGCTTTT	CAAAAAACAT	TATAAAATGA	AAAGCAGGAG	COTCACCAT
138181	GAGAATCACT	GGAACCCGGG	AGCTACAGGT	TGCAGTGAGC	CAAGATGGTG	CCACTCCACT
138241	CCAGCCTGGG	TGACAAAGTG	AGACTCTTCC	TAAAAAAAAA	ATAAATAAAT	A A TA A A TA A
138301	AAAAGAAAAA	GAAAAATCAC	AGGCTGAGAG	AAAATATTTA	TAATACATGT	ATTERESTAN
138361	GGACTCGCAC	CTGGAAAATA	TAAGGAACCT	TATAACTTAG	TAAGATGACA	ACCCARACAGA
138421	AAGAGTAAAA	GTTTTCAACA	GACATTTCAC	AAAAGAAAAC	ATACAAATGG	CCACTATCCA
138481	CATGAAAAGA	TTTTAAACAT	CATTAGTTAC	TAGGGAAATG	CAAGTCAAAA	CCAGIAIGCA
138541	ATACTTCACA	TTCAACAGAA	TAGCTAATGT	TABBACCACT	GACAATCCCC	ACCCRCAAIGAG
138601	AGGGTGTGGA	GGAAACTACT	CTCATATATT	GTGDATGTAA	GAGGACAATG	MUGGIGAGCA
138661	TTTGAAAAA	GTTTGGCTGT	TTCTAACATA	מסמממדדמממ	CTTATACAGC	CCACCAACIAC
138721	TTCTGGGTCA	TTTCTCCCAG	ATABATGAAC	ACATCTCCAT	ACTATGACAT	CCAGCAAIAI
138781	TCATACTGGC	TTTGTTTCAC	AATGCTATAA	ACTIGICAL	ACTAIGACAI	CCATCAAATGI
138841	GTGAATGGGT	AAATAAATTG	TAATATATCG	GCCAGACGCA	ACCCACGIGI GTGGTTCNTC	CCATCAACAG
138901	CAGAACTTTG	GGAGGCCAAG	ATGTACGGAT	CACCTGAGAT	GIGGIICHIG	ACACCACCCC
138961	ATCCAACATG	GTGAAACCCC	ATCTCTACTA	ALCOIGAGAI	TAGGAGIIIG	CACCCCCCCC
139021	TGTAATCCCA	GCTACTCGGA	AGGCTGAGGC	ADGDGDDTCA	CALCY VACCA	ACACCCCCAC
139081	GTTGCAGTGA	GCCAAGACCA	TGCCATTGCA	CTTCAGGGGGGG	CCCDDCDDCD	TGGN N CTCC
139141	ATCTCAAAAA	TAAAAAAAA	TGCAATATAT	CTATATOCCIG	CARCAMOA	AACCARDAAA
139201	AGGGAATAAA	CTACTGATAT	ATACACADAA	TGGDTGNATC	TCABARATCH TCABARATCH	CANCCAMIAAA
139261	AAAAATACA	TATGATATAA	ATTCCATTCA	TATCAMATC	TORMANIGI	AAAACTAACC
				GUMMIII	TUGGWITAGG	WWW TANGE

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139321	TGTAATTATG	GAAAGTACAT	CAGTGGCTGC	CTGGGGCCAA	GAGGATGGAA	GAGGCGGCAC
139381				TTGACGGAAG		
139441				CAAATTGTAT		
139501	ACTGTGTATA	AATTATTCCT	CAATAAAGTT	GATTTTTCAT	TAAATATATT	ATTTGCTAAA
139561	ATGAGGAGAG	ACAACTATTA	TCTTAAAATA	GTTAAGCACA	ATAAAAATAC	TACAATCAAC
139621	TCATTATATA	TGGAAATTAA	AGGAGAAAA	TAGTGGTATG	ATTAATTAAA	ATAAAAAGAA
139681	AACCTTCTAA	ATTTTATCTT	AGCTCATAGT	TGTAAAAGCT	GCCATCCCTA	ACCAAGGCCA
139741	CCCTTGACCC	TTTCTCATGT	TCCATCTTTC	TGTTTGTTTC	ATAGTTTATG	TCTCACCAAA
139801	ATCTATCAGA	TAAACGTATT	CATATGAAGA	TTTAAATATA	TTACATGTTA	AGCCTTAGCG
139861	AATACTTCAA	TATCTAAAGA	AGGTACAAAC	AAAACAAAAA	TCAACACTTA	GTTATAAGAG
139921	ATTACATACT	CTCCAGGGAA	GACCTGAAGA	CTAGCCCCTT	TCTGGATCCC	ACTAGCCCCT
139981	CATCCCACTC	CAAGCCCTCC	CCTCCAATCC	CATATGCACT	GGGCATTCAT	ACAAATAAGA
140041	CCATCAGCTC	TGGATATCTG	TACTGATTGA	TGCTCCTGCT	<b>AACTACCTGA</b>	ATGATTGCGA
	TGTAAGGACA	GCACTGCCTG	AATCCTATTT	ATCTCTCGCT	ATGCCATAGC	GGCCTTCCAT
140161				TCTTTGGTTG		
140221				GAAGAAATCA		
140281				AGTAGCCTTA		
140341				GGCTTTCATA		
140401	AGCTATACCC	CATATCATAC	ACAAAAGTTT	CTACATCTAA	CAAAGACACA	GATAGAAAAT
140461	GTTTTAAAAT	TTTAGAAGAA	AATAGTGCAG	AATTTTAGTG	CAGAATTTCT	TAGACTAGAT
140521	GCAAAAACAA	AAATGATTAA	AGTGGCCAGG	CACGGTGGCT	TATGCCTGTA	ATCTCAGCAC
140581				AGGTCATGAT		
140641				AAAAATTGGT		
140701	TTTAATCCCA	GCTACTTGGG	AGTCTGAGGC	AGGAGAATCA	CTTGAACCTG	GGAGGCAGAG
140761				CTCCAGCCTG		
140821				TTTTTAAAAG		
140881	TCGTTTAAAT	GATATGACAC	ACTACATCTA	ATATTTGGAA	AAGTACTTCT	TAATACTTTT
140941				TATCCTCAGA		
141001	GGGACGCAAG	CGCGTTCTTC	CTTCATTTTA	ACTGGTCATT	TTCATTTATT	TCAGGAACAT
141061	CTGAAGTAAA	CACAGTCACA	CGTTAACCTT	TAAAAATCTA	GGAGGTGCGT	ACGCATAGTT
141121	CCATTACTTC	AATTTTTGTA	CTTTTGCATT	TTAAAATATC	ACAGGGAAGC	TCGGTACAGC
141181	TTCAAGGCTA	GGAGGGGTGG	CTCTCTCTTA	AGCCCTGTCC	CCGCCAGCCC	CAGACCTCTC
141241				CACTTCCCCA		
141301				CTCAGACCTG		
141361				AAATTGAGAG		
141421	CTGTGGCTCT	CCAGAGGGCG	CGGCATTCAG	ACCCCAGATC	CAGCTGTGAG	AACGGACCCC
141481	AGGCTCACAC	CAGGCCTGCG	GGAGGCGGCC	CACCAGAGGC	GCTAGAAAAC	AAGCCTCGCG
141541	GGGAGGCGCG	CAGGGCGACT	GCAAGCTGTA	GGGGGCGCTG	GCGCCCTCAC	AGGCCAGGGG
141601	CAGGGCCGGC	GCTGCGGGCG	GGGCTCCTGC	GGCGTGAGGG	GCGGCCCCAG	GCCAGCAGCT
141661	GCGCCCTGGC	TGGGAGCCGG	GGAGCATTTG	CTGCTCTGCT	GGACCCTGAG	TCTGGCGGCG
141721	GGCGGCCTCC	TCTCCGCTCC	CCGCCCGCCA	TCCCCCAACT	CCCGATCTCT	CTGCTGCGTC
141781	TGGCCTCAGG	CTGAGACCCC	AACGAATCAT	TCCCCGCATG	GGAACATTTT	ATGATATAAC
141841	TGAATTCAGT	TTTATGTATA	ACTGAATTAC	GGATATGAGA	ATCTCAAATG	AGGACGAATG
141901	GITTITTACGC	ACAAAACATG	AGACACAAAT	CTGTAAGAAA	TATAAAGTCG	TGACCACGTC
141961	CTTTCAGAAC	TTTAACCTGT	TTGCTGAAGT	ACGTCAGTAA	CAATGGCAGG	GAAAGGGTAT
142021	CTTAAATTTC	ACCACAGCCT	CAAAGAGGCC	ATTTCGTGGA	TCCGCTGAGG	CTTGGAGTCG
142081	GCCTTCTGAC	CACGAGTCCT	GCGGCTATGA	AAGAGGAAGC	CGCGGTTCAG	GGCGTCCTCG
142141	TCCACATCGTGC	AGCCCGCCCT	GCTCCAGCTG	GGGACACCGG	TGGTCACGGC	GCTTTCCAGC
142201	CACCAGATECA	GGCGGCAGCC	CAAGATTTGG	TCCAGCCGCC	AAGGGGTGGC	TCGAGTGACT
142261	AACGGGCCTT	GAACGCTCCC	AGGACCCACA	TCTGGAGAGG	GAGGTGGGG	TGGGGTGCTG
142321 142381	AAGTCATTCT	ARCOMOCA	GGGGGGGC	ATGGACCTGG	GTAAGGCCAG	AGAAATTGAC
142381	ACCTCGTGAC	ATCUCTGGAA	GAGAAGTACG	TTCAGTGTCA	CTCCAGAGCT	GAAACCGCCT
	TOTAL	CCCTCCTCAC	CTACATACTT	TTCTAATTTG	TCTGGAGCAG	GCCGGGCATC
142501	TGTATTATCT	GGTTATTTAA	ATATCTGGTT	ATTTAAAAGC	TCTCCATTAA	ATTCACATAC

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142561						TAATGACCAA
142621		ATAGTGAATA				
142681		TAGACGCCCA				
142741		ATTCTTTAGC				
142801		CCTCATTTAG				
142861		ACTATTGCCC				
142921		CACTTTCACC				
142981	AAAGGTGAAA	ACGTTTCTTT	TATAATTTCA	CATACAATCT	TTAATGGACC	CAGTGTCCAA
143041		CAAGTGCTCA				
143101	GAGTTTAGGT	CTTGGAGAAA	AGAGACCCAA	GGAGACACAA	GACAAAGGGG	AAAGAGAAGG
143161		ACTGAGGACC				
143221	CGGAATATGA	CAGTTTGGAG	GGGCCTGAAG	GACTCTTCTA	TTCTCTATCA	GAAAAACAGA
143281		TAACCAGAAA				
143341	GTGATAATTT	AATGTGTTTT	AAAAAATGTA	TCACAGTGAT	GGCCTGGTGT	GAAATAAATA
143401	ATAAAATTTT	AAGAATTAAA	AAATATAAAA	ATCTTTTATA	TAGACATTAG	GAGTTACAAG
143461	GATAACTGTG	AATTATAATT	AGTAATTAAA	TTGAAATACT	GATTATTTTC	ATTTTTATTT
143521	AATTATTTAA	TAAAACCTAT	TTAACATTTA	ATATTTATCA	GTAATTAAAT	CTAATTGTTA
143581	ATATTTATTA	TTATAAATTA	TTTTAGAATT	AAAAATAAGT	GTAGAAGCGA	GGCATGGTGG
143641	CTCAAGCCTG	TAATCCCAAC	ACTTTGGGAG	GCTAAGGTGG	GAGGATTGCT	TGAGCCCAGT
143701	AGTTCLAGAC	CAGCCTGGGC	AACATGGAGA	AACCCTGTCT	CAATACAAAA	AAATGAGCCA
143761	TGTGTGGTGG	TGCGTGCCTG	TAGTCCCAGC	CATTCTGGAG	GCTGAGGTGG	GAGGATGACT
143821		CAGTCAAGGC				
143881		AGACCCTGTG				
143941		ATACAGAATT				
144001		GAGAGTGGAA				
144061		GCAGAACAAA				
144121		GGAGGATTGT				
144181		TCTAAAAAGG				
144241		TTTTCTGCAA				
144301	CAAAAATGAA	TAGATATTAG	TTGCCTGAAA	TAAAAATCAA	ATATCCAACA	AAAAATATTG
144361		AGTATCTAAG				
144421		GAAAACCATA				
144481		TTAGAAAACT				
144541		AAACAGTTGC				
144601		ACTTTGGAAG				
144661		ACTGTAGCAA				
144721		TTGCTAACAA				
144781		TTAGAGTACT				
144841		AAAGGACTTG				
144901		TTGCAAAAGA				
144961		CTCAACAGTA				
145021	TCTGTTCACC	AAAGAAGATA	CACAGATGCA	AGTATGCATA	TGAAAAGATG	CTTGACATCA
145081	TGTCATTAGG	GAACTGCAAA	TTAAAACAAG	TAGATACCAC	TGCATACCTA	GTAGAATGAC
145141	CAAAATTTAG	AACACTGTCA	GCACCAAAGG	TTGCAAAGAT	ATGTAGCAAT	AGTAACTTGT
145201	TCATTACTGG	TGAGAATGCA	AAATGTGCAA	TCACTTTGGA	AGACAGTTTG	GTGGTTTCTT
145261	ACAAAAGTAA	CCATACTTTT	ACCATAAGAT	TCACCAATCA	CACTCCTTAG	TATTTATCCA
145321	AAGGAATTGA	AAACTTATCT	CCACACAAAA	ACCTGCACAT	AGATGTTTAT	AGCAGCTTTA
145381	TTCATAATTT	ATCCAAAACT	TGGAAACAAG	ATGTCTTTCA	GTAGGTAAGT	GGATAACTGT
145441	GGTACTTCTG	AATAATGGAA	TGTTATTTAG	AGTTAAAAAG	AAATGCATTC	ACTTTGGGAG
145501	GCCGAAGTGG	GTGGATTGCT	TGAGGCCAGG	AGTTTGAGAC	CAGCCTGGTC	AACATGGGAA
145561	AACCCCAATT	AGCCGGGCAT	AGTGGCGTGA	GCCTGTAATC	CCAGCTACTC	GGGAGGCTGA
145621	GATATGAGAA	TCGTTTGAAC	CTGGGAGATG	GAGGTTGCAG	TGAGCCAGTG	CCACTGCACT
145681	TCAGCCTGGG	CAACAGAGCA	AGACTCCTCT	GTCTCAAAAA	AAAAAAAA	AAGAAAGAAA
145741	AGAAAAAAGA	AAAAGAAAA	GAAAAGAAAC	GATCAAGCCA	TGAAAACACA	TGAAGGAAAC

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145801	TTAAATGTAT	GTTACTAAAA	AGCCAACCTG	AAAAGACTGC	ATACTATATG	ACTCCAACTG
145861	ATGCAGGGCA	AGCAAGCCAA	AAATTAGGGC	TTAGCCCGGG	AAGAATTCAA	GGGTGAAGTG
145921	GTGGTGTTAG	CAACTTTTAC	TGAAGCAGCA	GTGTACAACA	GCAGAACAGG	TACTGCTCCT
145981	TGCTGAGCAG	GGCTAACCCA	TAAGTAATGT	GCCCAGAGTA	GCAGCTCAGG	GGCAGTTCTG
146041	CAGTAATATA	CCTGCTTTTA	GTTAAGTGCA	TGTTAAGGGG	GATTATGCAG	AAATTTCTAG
146101	AAAAAGAGTG	GTAACTTCGG	AGTAGGTACA	GAGGAAAGAA	GTCGATAATG	TCCTGTTGTT
146161	GCCATGGCAA	CGAAAAACTG	ACATGGCGCT	GGTGGGCGTG	TCTTATGGAG	AGGTGCTTTA
146221	ACCTCGTCCC	TGTTTCGGCT	AGTCTTCAAT	CTGGTCCGGA	GTAAAGTCCC	TGCCTCCGGA
146281					AGAAAAGACA	
146341	ACACAGTCAA	AAGATTAGTT	GATAGAAATT	GGGTGACAGG	AAGTGTTGAA	AAGGCAGAAC
146401	ACAGGATTTT	TAGGGCAGTG	AAACTTCTGT	GATACTATAA	TGGTGAATAC	ATGACATTAT
146461	ACATTTGTCA	AAACCCATAG	AAAGCACAAC	ACCAAGAATA	AACCCTAATG	TAAATTACAG
146521	ACTTTCGTTG	ATAATGACGT	GTCAATGTAA	GTTCAATTGT	AATAAATGTA	CTACTGTGGT
146581	GCTGGATGTC	TATGGTGGGG	GGACATTTTT	GCTTCAATAG	TTACAGTTGA	AGTAAATGTT
146641	TGTGTTTCCC	ACAATGCATA	TGTAGAAACT	CTCACATTCA	ATGTGATGGT	CTTTGGAGGT
146701	GGGCTCTTTG	GGTGATAGTT	AGGTTTAGTT	GAGATCCTAG	CAGATCGAGT	CTTCATGATG
146761					GCTAGCTCTC	
146821					AAGGGCCTTC	
146881					TTAAGTCACT	•
146941					TTAGAAATTT	
147001					AGCTTTTATA	
147061					AATAGTATGA	
147121					AATATGAGAA	
147181					TCTTAAGTGT	
147241			**		TAGGTTTTAG	
147301					TGTGTCCTTT	
147361					CTCTCTTGGA	
147421					TTTCCATACA	
147481					GCTTTAAATG	
147541					AACTTAATCT	
147601					AAAATCATCT	
147661	TTTGTTCCCC	ACATTATTTA	GTTGGAGCTC	TGTAACTTTT	TTTTTTTTT	TTTTTGAGAC
147721	AAGGTCTTGC	TCTGTCACTT	AGGCTGGAAT	TCAGTGGCAT	GACCATGGCT	CACTGCAGCC
147781					GAGTAGCTGA	
147841					GGGAGCCTTG	
147901					TCCCAAAGTG	
147961					AATTTCATTT	
148021	AGTCTGTTTA	GATTTTCTTT	CCCTTCTTGG	GTCAGTTAGG	CCATTGGTTT	CTTTTTAAAG
148081	GTTTTCAAAT	TTATTTGCAT	CTAATTCTTC	AAATTACTCT	CAAAATTATT	CCAGTATATA
148141	TTCTTTTGTT	CCTATTTTCT	TCTGTATTCT	TTATTAAAAT	AGCTAATGAT	TTATCTAGCA
148201	GGACTTATAT	TCTTTCCATA	ACTTTCCTGC	ACCCCAATTA	ATCTCCAATT	TTATATTTCT
148261	TCTGGCCTTC	CTTATAGTTT	CCACAGGTTT	ATTTTATTCA	TTTTTTAAAA	CTTTTATTTA
148321	ATTGTTTATT	TTATTATCAT	TCTTTCTTAT	TCAGCAATCT	AAGTGCTTAG	GGATATAGAA
148381	TTTCCTCTAA	GCAGCATATG	CTAGGCTTTA	ACAATGTTAG	GGAGGCCTCC	CCTTTCTGGG
148441	GAAGACCACA	CTTACATTAA	CACAGGACTG	TGGGATGCCA	AGAGGTAGAG	AAGAGCTTAT
148501					GGGGTTCCTC	
148561					ACGGGTATAT	
148621					TTTTCTATTC	
148681	CTCCATCTGT	GGCCCTATCA	AGTAGACTAA	CAAAAGACAG	ATTGACAAGA	CAGAAACAAA
148741	GCATGTGCAT	TGTACAAACA	CAGGGGAGTA	CTGAGATGAA	TACTCAAAAG	AGGATTTAGA
148801	ACTTGGGCTT	ATATAGCATT	TTAAGAAAAG	AATACATTTT	TTAAGTGACA	AGGAAGACGA
148861	AAAGGACTTT	GAGTTTCTAG	TGCAGTAAAT	TGTGGGAAGG	CAACTTTTTC	TTTCCCTTTT
148921	TTTTTTTTT	TTTTTAAAAA	AAAAGACTTC	TCTGGTGCTA	TGTCCAGGCT	GATAAGAGTC
148981	TAAAGTCTCT	GGTGACTAAC	TTTTGTTCTT	CCCCGAGTAA	GAAGACACCT	TCACAATTTC

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149041	ATATCCTGCT	TTTAGGCAAA	TAGGGAGAGG	GCAGAGGTGT	TTGTTTGTTT	TTAATCTATT
149101	TTTTTTCTCA	ATTGTCTTCA	ACTCAAAATA	CTTCTTATGC	CAAAGATGGC	ATATTCTGCT
149161	ACCCTTCACT	TACTACTTAC	AACCCAGCCT	CTATCATCAT	AATTAGAACT	TCTGACCCTG
149221		GCAATAGTTT				
149281	GCCATGCCTC	TGACATCTAG	ACACAACTGT	TGCTTCATTT	CTCCTATTCT	CAGAGGTGAT
149341	GTTGTAGGAC	TTCAACAAAT	ATCAGTAAAC	ATTAATTTT	TTTTTCCTTG	AGGCACAGCA
149401	TGATCTTGGC	TTACTGCAGC	TGCTGCAGGC	TCAAGCAATT	CTCCTGCCTT	GGCCTCACGA
149461	GTAGCTGGGT	TACAGGCCCC	TACCACCATG	CCCGGCTAAT	TTTTGTATTT	TTAGTAGAGA
149521	CAGGGTTTCA	CCATGTTGGC	CAGGCTGGTG	TTGAACTCCT	GACCTCAAGT	GATCCACCTG
149581	CCTCAGCCTC	ACATAGTTCT	GGGATTACAG	GCGTGAGCCA	CCATGCCTGG	CCATCAATTT
149641	TTATGTCAAC	TCTAAATTAT	AACATTTAGC	AATTTTGTGA	CTTTTTATGG	TCATCATTAA
149701	TGTTGTTTAT	GTTTTAGTTG	TAGTCCTGTC	ATTACTCACT	CGGGTATGGT	AATTTGGTCT
149761	TTTTCAAAAT	GAAGTTAAGG	TCTATTTGCT	CTTCTCTGAA	TCATAATAAG	AACTGCCAAC
149821	AGCCATTTCA	GCAATAACTA	TTTACTGAGA	TTTTAAAATA	TTTCAAGGTA	ATTGGTCCTA
149881	GCAGACTGGA	AAATACCAAA	TTCTTTTCCA	GAACTGAATC	CCCCATCAAA	GTTCAATTTT
149941	ACTCATAATT	CCCTTTTCAT	TTGAAGCATC	TCATTGTAAG	CCAGTCTTAA	CCCTTCTCTC
150001	ACACTTTGCT	TGGCTGTTTC	TCAGGTAGAA	CTCAGTAAGT	CTGGTAGCCT	CCAGGACTGC
150061	CGCTTAGATT	ATTAAACAAC	ATGTCAGTGG	TTGGAAGAGT	CAATGTTATT	TTGATTTTTC
150121		TGTTTTAAAT				
150181		TGGCAGCAAA				
150241		CCTGAAGACC				
150301		ACGATCATCA				
150361		GTTGCTAAGT				
150421		GTCTAACTAC				
150481		TTATATTATT				
150541		TGGGGGAGGC				
150601		AGAGAAGGAA				
150661		TACAACTCCG				
150721		AAGGGAAGTT				
150781		CCAGAATCCA				
150841		AACTCCTAAA	4			
150901		GGTGAAGCCT				
150961		CTGATTCTCC				
151021		GTGATGAGCT				
151081		AAATAATTT				
151141		GTGTTGCTGT				
151201		CTCCTGGATT				
151261		CACCACCACA				
151321		GCCAGGCTGG				
151381		CTGGATTACA				
151441		CTGTGTTCAA				
151501		CTTTCTGAGC				
151561		GGTGGGTAGG				
151621		GTAACCACCT				
151681		CTACAAAGTC				
151741		GTCCAGGGTG				
151801		TATTTATTGG				
151861		GCATGAGGAC				
151921		ATTTTCTTTG				
151981		GAGTGAAAAG				
152041		GCCCTGAGCC				
152101		TGGTGCGGCA				
152161		GGGGTTTTGG				
152221		GCCAGTCCTG				

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152281	CAACATCATO	TTCGTCTTAA	TTATTCAAGG	ATGCCAAGG	ר מרמקממרדמי	A CCTGTTAATA
152341	TGGTTACCAT	r cctgtccaaa	GTTCTTCTCC	CATGCAGGA	TTCCAGGAAT	CATCACACAC
152401	TTGAGCAGA	AGATACCTTT	TCCCTTCTCT	ACTGAATAA	CACCAACATT	GAGAATCAGA
152461	GAGGGAAAAT	GACTCAGCTA	ATGTCTTAGC	TTGTTATTG	A A G A C C C A G G	TCTCATGACA
152521	CATGCCTAGI	CCCATGACTT	TTAATTGTAA	GCTCTTCTC	TTCCCCTCAC	ATAATGTTCC
152581	ATAAGCATTA	GTATGAGATA	ATAATACACT	GAGGACCAA	י בובכככנונאט	ATATIGTICS AATATCAGAC
152641	TAGAATCAAA	CAAGACAGAA	AAAAGATCTG	ב מדשם כרכים	CTCACATOAAA	GAACAGTATG
152701	CAGTTTTAAA	TAAAAAAAA	GGTAATAGGA	TGTTCTAAC	ACACACTURA	GAACAGTATG
152761	TGCTACTGAG	TTAAATGTTG	ATCAGTTGGT	CTGTGACAA	TANCCARTAG	AAGTATTCAG
152821	AAACACTTCC	TGTGCTGGAT	GCTCTCTGTT	TGTTCTTCC	ANTANTOCOM	CACTTTTCCC
152881	TGTCTTGCTC	TGTGCCCAGG	AAGGCTGACA	TGGACAGATT	, WILWILCCCI	CACTTTTCCC
152941	GGCTTGGTTC	AGCCAATGGG	AAGCACCAGA	GGAGACCATA	- AACCAGGCIT	AAGCAGCCTCT
153001	GGGAGTATTC	AGTACCCCAG	TCCCACGCTA	TCATTTCCAC	CCTCTCCA	CCTCTGCCTC
153061	TGGGCACACT	CTAGTATAGT	TACAGCTCCC	TACACCTCCC	' ACTTCACATT	CAGAGGAGGT
153121	GATGGCTCTC	TAACTGTTCC	TAGTTCTGGG	TACACCIGCC	. ACTIGAGGCC	CAGAGGAGGT TTTCCCAACT
153181	CCTCACCTTT	GTANATACCC	TAGIICIGGG	ANGREEN	TCCTTGTGGA	TATCAGCCTG
153241	ACTCACAGAA	GTTTGGGGTT	TCCITITICA	MACICIATIC	AGTTAGCTTT	TATCAGCCTG AACCCATGTT
153301	GAGAAATTAA	AATGTTTACG	GGGTGGTAAT	1 IACCTGAAT	GACCCAGGAA	TCAATTGGAT
153361	TTTTAAAATT	CCACCTATCT	ATTCCTCTCA	CACACTTAAG	AGAAAAAATA	TCAATTGGAT GAAAGATTGG
153421	AAGCTAAAAG	ATAGATAATA	TACTCATATA	CACATCAACA	AAAACATATA	GAAAGATTGG AAGATATTAA
153481	GTCAGAGCAT	TATTAAGAAT	GGAAGAACCC	COLCOMORGO	ATTATATCAA	AAGATATTAA
153541	AGCACTTTGG	GAGGCCAAGG	CAGGGGGAAGGG	CCAGGTGTGG	TGGCTCATGC	CTGTAATCCC
153601	CCCAACATGG	CAAAACCCTG	CCTCTACCAA	ACTIGAAGCC	AGGAGTTCAA	GACCAGCCTG
153661	CATGCCTGTA	ATCCCAGCTA	CTTCCCACCAC	MAATACAACA	ATTAGCTGGG	CATTGTGGCA
153721	GCAGAGGTTG	ATCCCAGCTA	CIIGGGAGGC	TGAAGCACAA	GAATCACTTG	AACCGGGGAG
153781	ATTCTGTCTC	CAGTGAGCTG	AGATTTUGUU	ACTACACTAC	AGCCTGGGTG	ACAGAGAGAG
153841	CAATTTTAAA	AAAAAAAAA	CTACATEATE	AATGAAAGGA	GTCACCTAAA	AAAGATAACA
153901	TACADAGCAA	CATAAATGTA	ATTEMPORA	AGTGAATTCA	TGTTTAGAAT	TGTGTTAATA
153961	GTTTTAACAT	AAATTGTAGA	ATTATAGGAG	AAATGGACAA	ATCTACAATC	ATCATGGGAT
154021	GAAGATACGG	TCTTCTTTCC AAGGTCTGAA	CARTCTARCA	GATCAGGCAG	ACCAAAAGAA	AGAAATAAGG
154081	CAGCAATTGT	TTAATAATAG	TARCACACA	AGCGCAATCT	CATAGTCAAT	ACATAAAGCT
154141	ATGCACTAAC	TGAGTAAATA	CTAGGGAGA	ATATGCAGTT	TTCTCAGGTA	TAGATGGAAC
154201	ATTACACAGA	TCATTTTCTC	TAGGGGGAA	AACAGICIGA	ACAAGTTTCA	ATAAATCTGT
154261	ACTAAAAAGA	TTCTAAATAT	TAGCCICAAI	ALAGATTAT	AAACCAATAA	TAAAAAGATG
154321	AGAATGGAAC	TTCTAAATAT AATAATAAAA	ACTTATOTAT	AAACTACTAA	TAAGTCATTA	GAAGATGTAT
154381	TTAAGGAAAA	TTTGTAGGCT	TTARATION	AAAAATATAC	AATGAAGCTA	AAGCAGAATT
154441	ATGAGCCAAG	CATCTAATTT	TIMMIGCII	AICTIAGAAA	AATTAAAAAG	CTGAACATTA
154501	AAAAAGAAAA	CATCTAATTT AATAGATATT	AMMIIIIAMA BBBCBBCBCBCBCBCBCBCBCBCBCBCBCBCBCBCBCB	AAGAACATAG	AAAGCCAAAT	ATAATTTTTT
154561	TAAAGAGGAA	AAACAAACAA	AUTHORNIA IN	ACAGTGAAGT	TAAAGAAAAC	AAGAATGCAA
154621	GACATACCTC	CAATGAGATT	TATCA A A CTA	AGCTTCTTTT	AAAAGAAATT	TAATAAAATA
154681	AAACTTTTTA	AATATTACAG	TATCAMAGIA	TA A A TOTAL	CACAAATGGA	ATGAATACAG
154741	TACTGATAAA	ATTATTACTT	CCTACAAAAA	ATATOTTAT	GCTACTAATA	AAATTGAAAG
154801	AAATAAAGCA	ATTATTACTT TGGGCAGACC	TARCAMANA	CARACTECTGA	GTAAAACTCA	CTCAAAAAAC
154861	ACAGATAATA	AAACGTGCAT	CTTTTTTTTTTTTT	GAAATGAAAT	CACTACTITA	AATTTTACCG
154921	ATTTAGAAGT	AAACGTGCAT CAAGGCATGA	CTITALCAAG	CAAAAATGGA	ACTTGTCAGT	TTTATAGGAA
154981	ATTATGGCTC	CCGCTTATAG	OTANIOCCAM	ACTACCA	AATCCTACAA	AGAATAGAAA
155041	AAACCAAATT	TTATATTTCC	77CTAGA1A1	AGRACICCIG	CACAAAATAA	TATAAATAAC
155101	ATATACATAT	ATAATATGTA	TRECRETACE!	ATTATATGTG	TATGTATTAT	ATATGTTAAC
155161	ATATACATAT TTTTACAATA	TATAAATGAA	PUCCUINIGI	TTATAUATATT	ATATATGTAT	AGTGTATGTA
155221	ACATATATAA	TACATTACAT	CARACAMICI	TACAAMAATATT	CATCTAGATT	GTCATATATG
155281	TGCCTGTAAT	CCCAGCACGT	TCCCACCCTC	ACCCCCCCCC	AGGCCAGGCA	CAGTGACTCA
155341	TGAGACCAGC	CTGGTCAATA	TCCCCD B ROOM	CCATCTCTA	ATCACTTGAG	TCCAAGAGTT
155401	CAGGCATTGT	GGTGCACACC	A DECEMBAL I	CCALCICTAC	AAAAAATATG .	AAAAATTATC
155461	CTTGAGCCTG	GGAGGTGGAG	ATTICITUM	GCIACICGG GTCCACATTTC	AAGCTGAGGT	GAGAGGATCA
			WITGCWGIGW	GICGAGATTG	CGCCAGTGCA	CTCCAGCCTG

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155521		GGAGACCCTG				
155581		GTCCCAGCAA				
155641		GATCGCACCA				
155701		AATATATACA				
155761		TATATAATAT				
155821		TACAGATATA				
155881		AGCATATAGA				
155941		CTGTAGTCCC				
156001		TTGAGCCATA				
156061	ACCTGAGGTG	GAAGGATATA	GATATAGATA	TATAAATAAA	TATGTATAGA	GAGAATATAA
156121		TATGTGTATA				
156181		GTGTATATAT				
156241		TCCAGGTATG				
156301	CACTGGTCAT	AGTTACGGGA	AAAGAAGGTC	TCCAATGAGA	CATACTTAAC	AAAATATATG
156361	AACTTGCCAT	ATACGTGGAG	AGTTCTGGTG	TGTATATAGC	CTTCTCTCAC	CAACCTAGCA
156421		TCATCATTAT				
156481	CCTTTCTTCT	TCTTTCTCTT	CCTTCCCCTC	CCCCACCTCT	TTCTCTTCCT	CCTCCTCCTT
156541		CTTTTTTTTT				
156601	GTGGCACAAT	CTCAGCTCAC	TGCAACCTCT	GCCTTCTGGG	TTCAAGCAAT	TCTGCCTAAG
156661		AGCTAGGACT				
156721	TAGTAGAGAT	AGGGTTTCAC	AATGCTGGCC	AGGCTGGTCT	CAAACTCCTG	CCCTCAAGTG
156781	ATCCTCCTGC	CTCGGCCTCC	CAATGTGCTG	GGATTACAGG	CGTAAGCCAC	TGTACCCGGC
156841	CTCCTCCTTT	AATAGACAGG	GTCTAGCTCT	GTTGCCCAGG	CTGGGTACAG	TGGCGTGATC
156901	ATAGCTTACT	GCAGCCTCGA	ACTCCTGGGC	TCAGGAGATC	CTCCTGCCCT	AGTCTCCCCA
156961	GTAGCTGGAA	CTACAGGCAT	AGCACACGGG	GCTAATAAAA	TTAATTAGGT	GATAAAATTC
157021	ACTGCCCACT	GATGACTAAG	CTCTTTGGAC	ATAAAAGACA	CAGACCTTGA	AGGAAAATGT
157081	GTCTACTTAA	TTTTGAAACC	CTATTTATCA	AAAAACAGGA	TGAAAATGCA	AAATGCCATC
157141		AAGATATCAG				
157201		TTAAACCACA				
157261		AAGAATCTCA				
157321		CATTITAAAT				
157381		TTGGAGGAGT				
157441		GTAGTATCTG				
157501		TCCCTGGAGG				
157561		TTGTCTGCAA				
157621		CACAAGGCAA				
157681		AACACTTTAG				
157741		AACAAAATAA				
157801		GTGCTTGCCT				
157861		GAATTTAAGG				
157921	GTGACAGAGT	GAGACCCTGT	CTAAAAGAGA	TAAGTAAATA	ACAACTTTGC	ATTTTCTGCC
157981	ACATTGCAAA	ATGGTGAGAG	AGTGGTTTCT	AGACTCTAGA	CTCTTTCTAT	GACTACCTTC
158041		ATCCTACAAC				
158101		CCCCATATAG				
158161	CTTTTCTAAT	CTGTCACAGA	CTAAAGAGTG	CTCAGTATAT	GTGAGTCATT	ATTCCTGGTG
158221	CIGGIAGGAG	TGTATGTTAC	AACTITGAGT	CAAGTAATAT	GGTACCATAT	ATTAAGATTA
158281 158341		CGGCAATCCC				
158401		CATGGACTAG				
158461		GCTCCATCAG				
158521		AAAAAGTAAC				
158521		GGAAAGATAT				
158641	GAMMACCITI	GAACTTTCTC	CITATATCCT	TTATATTGTT	TGACTGATTA	AAATGTATTT
158701	DITECRICIC	CTTGAAGGCA	AIGTAAAATA	AAATAAACAT	ACATTTAAAA	ATAAAAATAA
20.01	WITTIMITCO	TATCACTTTT	GTAATAAAGC	TGGGCACAGT	GACTAACACT	TGTAATCCTA

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158761	GCACTTTGGG	AGGCAGAGAC	AGGCAGATCA	CCTGAGGTCA	GGGGTTTGAG	ACCAGCCTGG
158821	CCAACATTGT	GAAACCCCAT	CTCTACTAAA	AATACAAAAA	TCAGCCAGGC	ATAGTGGTGC
158881	GTACCTGTAA	TCCCACGCTA	CCCGGGAGGC	TGAGGCGCTG	GAACCCAGGA	GGCAGAGGCT
158941	GCAGTGAGCT	GAGATTGCGG	CACTGCAAGC	CAGCCTGGGT	AACAGCGAGA	CTCCATCTCA
159001	AAAAAAAATT	TGAAAAAAGA	AAAATTTTAA	TAAACAGTGT	TTAAGAGGGG	AGAAATATTT
159061	AGTTAAAAGA	TAAGCCCATT	TAAGAAATAG	TTTCACTTGA	CCCGGAAGGC	GGAGCTTGCA
159121					AGAGCGAGAC	
159181					ACTTGAACCA	
159241					AATCCCAGCA	
159301					ATACAGAGTG	
159361					TCTATAATTC	
159421					AGAAGTTATA	
159481					GCAGAGCCCT	
159541					TGGGATAAAT	
159601					GGAAGACAGA	
159661					AAAATTCCTT	
159721					GGCAAAGATG	
159781					CTGGCAGATT	
159841					TGTTAGAAGC	
159901					AGCCAACAGC	
159961					CTGCTACCTC	
160021					CAGCCTGACC	
160081					CTGTGCCCAG	
160141					CTAAATTATG	
160201					GGTTAGGCCA	
160261					TTTTTGATTT	
160321					AATCAAGGGA	
160381					GTAACATTTG	
160441					TCATACTCAC	
160501					ACTATGATCA	
160561				*	AGCACTTTGT	
160621					TAATATTTTC	
160681					GAAAGAATAA	
160741					ACTGAGCAGT	
160801					CAACCTCTCT	
160861					CCCCTTCTAT	
160921					ATTTTCTTCA	
160981					ACAGCCTCTG	
161041					TTCTTCAGAG	
161101					AACAGGACTT	
161161					GAGCATGGGA	
161221	TTCATGGGGC	TTGTCATGCA	GGGCTATTCT	TCTTTCCTTT	ACCCGAAGAA	GTANAGAGAG
161281	TTACCCTAGT	CTTAGTCTTA	GATATTGATG	GATACTCAAA	CAAAGTAATT	CCCACCAGTC
161341					GCTTCTGGGA	
161401					GCTGAAGTCT	
161461					GATCGGATTT	
161521					GGTGAATCTG	
161581					TCACACAGTG	
161641					GTTAGTTAGA	
161701					ATTGTACCAT	
161761					CTCATTTTGG	
161821					AGTCAACAAC	
161881	TGCACATGGG	CCAAGGAGGC	AAACAGTGGT	AAATGTTATC	CCGTGGTTTC	ATTTGGCCAA
161941	GCTGTGTTCC	CTCAGAAGTT	TATTTTTCTA	ATTGACATAA	AGGTACCCTA	TAAATTAGTG
_					A	**************************************

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162001					ATTACTTTAT	
162061					TTTTCCCTAC	
162121					TTTTCCTCCA	
162181					TCATAGTCCC	
162241	CAGAGTCTGT	TGTTCTTCTT	TCTCCAGCCT	CCAGAGATAA	GACTTCTCTT	CCTCATGTAG
162301					CTCTGAATCT	
162361	CTGGAGTCAA	GAAAGTATGG	TCAAAAGGTG	GAAGTAAACC	AAATGTCCAT	CTATGGATGA
162421	ATGGATAAAC	AAGAATGAAA	GTCTGACACA	CGCTACTACA	TGACAAGCCT	TGAAGACATT
162481	CAAGCAAAAT	AAGCCAGAAA	CAAAAGGGCA	AATATTGTAA	GACTTTGCTT	ATACAAGGCA
162541	TCTGGAGTAG	TTAAGTTCAT	AGAGACAGAA	AGTAAAATAG	TGGTTACAAG	GTGTTGGCAA
162601	GACCAGAAAA	TGGACAGTTA	TTGTTTAATG	GGTAGTGAGT	TTCAGTTTAG	AAGATGAAAG
162661	ATGAAACTGA	GTTGCAGTTT	GGAGATGGGA	ATGGTGATGG	TTGCACAACA	ATGTAACAAT
162721	GTAAAAGCAC	TTAATTCTAC	TGAACTATAT	ACTTAAAAGT	GGTTAAATGC	TTAAGTGTTA
162781	TATATATTTT	CACACAAACA	CACACACACA	CACAATCAGC	CACTGGGACA	TTATTTTCTC
162841	ATGAGTCACT	GAAGCTGGAA	GAATGTCCCC	AGTTTCCTGC	TGCAGAGTCA	TGTGTGGGAG
162901	GCAGGCACTC	AGATGTGGAA	GAGGTTGCCT	CAGATTCCTT	ATAGTCACCC	AATTAATTTT
162961	CTTGTTCTTC	AGCCAAGACA	CAGGAGAAAG	CTGGGTTAGG	AGTGCTAGAT	AATTTAATTG
163021	TGAAACTAGG	GCCAAGTTCA	AACACTTTAT	CAGTTACAAG	GATAAAAAGA	GGTTTTTACT
1630B1	TATGATTTAA	GAAGTTAGAT	TTCTGAGTTG	GAGCGATTTT	CTTGAAGTAA	AAGCTTATAA
163141	TGAACA'TCAC	CCAGACTGGA	TTTTAAGACA	ACCAGGCTGG	TAAGAGGGTC	CATAATTCTT
163201	GGCAGGGGGA	GCTTTGAGTG	TGACAGGCAT	TTATTATGGT	TAACTGAGAA	ATACTGTTCT
163261	ACTACCCTAG	GGTCATCTTA	AGCATTCCTA	TGTGTAAGAC	TGACAGAAAT	CAAGTGAAAC
163321	TCTCATCTGA	GGAGATGTAA	AGTTGCAATT	TCCATTAGTG	CTGTCTAAAT	TAATGCAGTG
163381	GGAGTGTGTA	TTCAGGGCAA	TTTGAATCTA	TGTTCTTGGA	TTGCAGTCTT	CAAACTTGGC
163441	CCAAATAAAC	TCTCTACTTA	TCTTAAAAAA	ATAAAAATTA	AAAAATAAAA	ATAAATTCAT
163501	ACAGTGTTTT	GATGACTATG	ATATAGAAGA	AGGGTCTTTG	ACTTAGGATG	AGGTGGAATT
163561	TTTGTGTAGG	AGACAGGTGC	AGCTTTAACT	CTTGTATAGA	CGGGTTTTCA	TATATGTTAG
163621	TTACAATCAA	GGTCTTCCCC	ATTGCCCAAG	ATCCTAGAAA	TGGGGGAAGT	AAGAGTGTAC
163681	TCAGGAGCTC	AAGAGCAACA	TCCACAAACA	AAGATCAGGG	TAGAGGTTAG	AGAGGACTCC
163741	TGAAAGAGAG	AAAATTGGTA	ATCAGCTTGT	GGGATTTTAC	TGCAAGCTAG	TGAATTATAT
163801	AAATATAAAG	ATTGGTGCAA	AAGTAATTGT	GGTTTTTGCC	TTTACTTTAA	TGGCAAAGAC
163861	CGCAATTACT	TTTGCACAAA	CCTAAATATT	TCCATAAAAG	AATGTGGCTC	TGATAATGTG
163921	GAGGTTAGTC	AGCCACGGAA	ATAATCTGAA	AGTTTGTAGT	TGCAAGTGTG	TAGGTTGTTG
163981	CATTACTTGT	GATGTACTTA	TAAATCAAGT	ATAGGCCGGG	TGCAGTGGCT	CACGCCTGTA
164041	ATCCCAGCAC	TTTGGGAGGC	TGAGGTGGGT	GAATCACGAG	GTCAGGAGAT	CAAGACCATC
164101	CTGGCCAACA	TGGTGAAACC	CCGTCTCTAC	TAAAATACAA	AAAATTAGCC	AGGCATGGTA
164161	GCACATGCCT	GTAATCCCAG	CTACTCAAGA	GGCTGAGGCA	GGGGAATTGC	TTGAACCCGG
164221	GAGGTGGACA	TTGCAGTGAG	CTGAGATCGC	ACCACTACAC	TCCAGCAAGA	CTCCATCTCA
164281	AAAAATAGTA	ATAATTTAAA	AATAAATAAA	TAAATAAAGT	ATATTTCTTT	CATCAGCTTC
164341	ATGAGCTAGA	GTAGTATGAA	TTTCAATCTG	GAGTGATCCT	GTTTTCTAAG	TGTTCACAAA
164401	GCTTGGTTTC	TGTACCTGTA	AAGTTGAGAG	CCAGATGCTC	CACTGTGGTA	AAAGTGCCAG
164461	GGTAATGAGT	TGAGGCCTGC	AAACCAGGTT	TATTTTGACG	TATTTAAAGT	TTGAGACCCA
164521					TTCTTCTGAC	
164581	GGAATCCCAG	CCAACTACAG	TTTAAAGATG	GAAAGATTGG	TGCTAAATAC	TCATGGATGT
164641	AAACCTGGAA	CCAGGGGCAT	AAGTACAAAT	AATGGTTTCT	TCCTTGGGTT	TCATTTTTTC
164701	AATCTGGTTT	AGTGAGAATA	AATCCTCATT	GTGCTTTTCC	TCAATCATCC	CCTATGCCTA
164761					ATTCTTACTT	
164821	TATTCGCATT	GCTGTGGACA	GCTTCTGCTC	CGTACATCTG	TCTTCAAGTT	GCTTCAGTTT
164881					ATAAGTGAAG	
164941					TAAGATATAT	
165001					CATGCTAAAG	
165061					TAGGATTTTA	
165121					CTTGCCCAAA	
165181					CCCAGAGCCT	

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165241						TCATCTAAAA
165301					CCAGTGTTTG	
165361					CTGGTTATTA	
165421					CATGAATCCA	
165481					CAGTTTCTCA	
165541	GAAAGTGATT	TGAAGCTGAC	CCAAATTGCT	AATTGTAGTC	AATGCTGAAA	GAATTGTCTC
165601					GAGTGTTCTC	
165661	TCTATGTAAC	TGTTTTAGCA	AAAGATGACA	TTGTCCTTAC	TATATGCCAA	GTGCTATTCT
165721	ATGCATTCTA	TATTTTAATG	TCCTCAAAGC	TTATAACCAC	CTCCTGTGTA	TGTGTTTTAG
165781	GGAGGGAGGA	CACTGCTATT	ATCCCCATTT	ACAGATGGAG	AAACCAAGGT	GTGAAGACAT
165841	TAAGTAACGT	GCCCAAAATT	GCCCATCTAG	TAAGTGACAA	AACTCAATTT	CAACATAAGC
165901	TGGTTCCTTT	TCTTACTACT	TGGTGGAAAA	GTAATTCAAA	TGGGAATATG	ATCATCGCAG
165961	TTATTAGCTG	CTCCATGGAG	TTTAAGGAAG	AGCTGCCATG	AGCTGAGTGG	TGGTCATGAT
166021	TGACATGTCC	TTAGAAGGAC	TTAGAGCCTT	CATACAAGAC	CACCTCTGCC	TCATGGAGGA
166081	CAGAATAAGG	AGCCTGACAC	TGGAGACAAC	ATTTTCCTCA	AATTTAGGCA	GGACAGAGAA
166141	GGAAAAAGGA	CATCAGGACT	ATGCCCATTC	CTCCATGCTG	CCAACAGCAA	AGTCCCACCT
166201	TCCTTAATAT	GCTTTCTGGC	AAGAAATCTG	GATGGTACAC	AAAACCTCTC	CCTCTGCTTC
166261	ACCTTCCACA	ACCAAGCATT	TCCAAATCTT	TGACTCTTCT	TCCTGAATCG	TGCTTAAAAT
166321					TACTCCTTGA	
166381					CCTCTAATCC	
166441	GGAGGCTGAG	ATGGGAGGGA	GACCAGGGGT	TTGAGGCCAG	TATAAGCAAG	AAAGGCAGAC
166501	CATGTCTCTA	CAAAAAATAA	AAAAATTATC	CAGGTATGGT	GGGGCATCCC	TGTAGTCCTA
166561					AGAAGGTTGA	
166621	AGCCGAGATT	GCACCATTGT	ACTCCAACCT	GGGATACAGA	GCAAGACCCT	ACCTCAGGAA
166681					GATATTTTCA	
166741	CCCTTCATTC	CCCAAATGAA	AATCCCCCAA	TAGGTGTTCA	ATTTTTACGT	GTCCTTCAGG
166801					CTCCCCACCA	
166861					CTAGACTGTA	
166921	AGGTGATGGG	TCTTTCTTCC	CTGTTTTCAG	GCCCTACTGC	ATGGCTTTAC	ATATTGTGGT
166981					GGGTGTTTGG	
167041	ACCTGTTCTA	AAGCAAAAAG	AAATTCATCA	TAACACAAAT	GGATAGAGAT	AAGAGTCCAA
167101					AGCAAGAAAT	
167161	TATTTTTCAG	AAGAATGACA	TGATGAAAGC	TGTATTTCCA	AGTCATAATG	TTAGGTTTCA
167221					GGTACTTACC	
167281	GGCCCACACA	CTCACCTTGT	AGCCCTGGCA	TACGTCTTCA	ACAAGAGCTG	TGGTGTGCCC
167341					TGCCCCTCGT	
167401	CAGGTGGAAC	TGCTCTCCGT	GTTCCTCACA	TGACATTTCT	TGATCCGTCT	CTTTGAGGGC
167461					TCCATATGAA	
167521	ACACTGGGGA	CAGCAGAATG	TCTCCTGCCT	CAGTTGCTTT	TGGCTTGGGT	TTTTAAAGAA
167581	GTCTGTTATA	CACAAGTGGC	AGTAGCTGTG	TCCACAGTTG	ATGCTTACTG	GGTTCGTCAT
167641	CAGGCTCAGG	CAGATGGAGC	AGGTGGCTTC	CTCCATCATC	TTCTTGGTGC	TGGTGGTTGA
167701					AGATGGAGAT	
167761	GAATTTTCCA	CCGTGATGAA	AATACACCTC	ACCTGCACCT	CTATGTGATG	AGCTGGCTGC
167821	AACTGACTTC	CATAGGTCTT	GAAGGTTTTC	CTTCCAACCC	CTATTATCTC	ATTTTGTATT
167881	GAAGAAAAGA	GGACCTAAAA	GGAAGAAGTT	GAGGCTGAGG	TTGTTTGGGC	CACGTTTGAG
167941					GCAAGCAGTT	
168001					TGTTTGCCAA	
168061					TGTATTACAA	
168121					GCTTTTAAAC	
168181					TTTTGCATAC	
168241	TCAGACTGCT	CTAAATTATT	TCATTATTTT	TCTTTTCTCA	GTCTTCTAAC	TTTTTTTTT
168301					GAGTGCAGTG	
168361					CTGCCTCAGC	
168421	GTAGCTGGGT	CTACAGGTGT	GCACCACTAC	GCCCAGCTAA	TTTTTGTATT	TTTAGTAGAG

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ATGGGGTTTC ACCATGTTGG TTGGCTCGAT CTCTTGACCT TGTGATCCAC CCGCCTCAGC
168481
168541
       CTCCCAAAGT GCCAGGATTA CAGGCATGAG CCACCGTGCC CAGCCTCTTT TTCTTTCTT
168601 ATAAGACAAG TTCTCGCTCT CTTGCCCAGG CTGTAGTGGA GGGCAGTGGC ATGACCACAG
168661 CTCACTGCAG CCTCGACCTC CTGGGTTTAA GCAATCCTCC TGCCTCACCC TGGCAGAGTG
168721 GCTGGGACTA CAGGTATGTG CCACCATGTC CAGCTAAAGT CTTCTCCCA GAAAGAAGAA
168781 ATGCATTGGA ATTTAGAGGA TACACAAACA TCTAGCTGTA TAGCTAATAC AGTAGCCACT
168841 ATCATGAGTA GGAATTTAAA TITAACTTAA TAAAAATTAA AATGAAAAAA TTCAGTTTTT
       CTGTTCCAGT TGCCACATTT TGATTGCTTA ATAGTTGCAT GTGACTAGTG GCTACATAAC
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168961 AGCCTCAATA TACAACATTC TGTTATCACA GAAAGTTACC TTGGACCAAG TGCTGGGAGA
169021 AGCAATGCAG GCTTCCTCAC AAAAGCTGTA AAAGAGAGAA CTCAGGGAGT GTGAAACTCT
       TTCCTATTCT AGTTAACTTC AAGAATAATT GTTACCAGGC CAGCACGGTG GCTCACGCCT
169081
169141
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       CCAGCCTGAC CAACATGGCA AAACCTCATC TCTACTAAAA ATACAAAAAG TTAGCTAGAT
169201
169261
       GTGGTGGTGC ACACCTGTAA TCCCAGCTGC TCAGGAGGCT GAGGAAGGAG AATGACTTGA
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       GCTCCGGAGG GGGAGGTTGC AGTGAGCCCA GATTACACCA CTGCACTCCA GCCTGGGTGA
169381 AAGAGCGAGA ATCTGTCTTA AAAAAAAAA AAAGAATAAT TGGTACCAGA ATTACTCTTT
       GTAATTAGTA GTAACACTTA TGCAATTGGG TGATCTGTGA CAGATTCCAT TGAAGGAGTA
169441
169501
       TGGGGAGCTT CACCCCAATA TATGACTCCC TGGTATAATG AGTATTTTGA ATTAAAGGCC
169561 CTTAGAGATC AGCAGATGCT GGAAGAGACT TTTCCCCTAT CTACATAAAG ACCAGTCACA
169621 CTAGACAAGA AGAACAATTG TTTTTCCTTC CAACCCCTAT TATCTCATTT TGTACTGAAG
       AAAAGAGGAC TAAGAATGTA ACCAGACCTA ATCAGACACT TTCACAAAAT AATGTCTGTC
169681
       TCTCAGGCTC ATTCATTTTC CAAAGAGAAC CATTTACAAG TTAAACTCTG TTCCTCCATT
169741
       CATTCATCCT CCCAAATATT CATTTATTCT CCCTAGTAAT CATTTACTGC CCCTCAAAGA
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169861
       ATTACCTATA TTCTCCTGAT ATCACCCTTC CCCTCTGAAA TAAATATGTA TACATGTATA
       AACGTTATAC ATACATATTT ATACAGTATA CATACATATT TATACATACA TACATATGCA
169921
169981
       TACATATTTA TATTTATGTA TTTATACATA AGTATTTATA AATAAGGCTA TATAAGTATC
       TACCCCCATT GGCAGAGGGG GTAATCACTC TGTGATTCTA GCCCATGTAC TTGTTAATAA
170041
       ATTTGTATGC CTTTTCTCCA ATTAGCCTGC CTTTTGTGAG TCGATTTTTC AGTGAACTTC
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170161
       AGAAGGCAAA GGGGAAGTGT TCCCTTGGCT CCTACACCAT CATGACAATA AAATTTGACT
       CCACCTCGAC CCCCCCATC CCCCACAAAG AACAACAACC AACACTGGTT AATAAGGTCG
170221
       GTTGTTTTT GTTTGTGTTT TTGTTGTTGT TGTTTTTGCT TTCAGGAGCA GAGGTATAAT
170281
        AGGCAAAAGA AAGAGAAAGG AGAATAGTGA ATACCTCTTC TGCAGAGAGG GGTGCCTAAG
170341
       TGGGACTTCC CTGGCTAATA ACGTCTTGCT AGAGACCCAA CCAGGAGGAT AATGGAAGCA
170401
       ATCAAGGCAA CCAGAACAAC CAGAAGAACC GGTTTATCCT TTTTGTGCCC TCTCCCTAAA
170461
       170521
       TTATTTCTAT GGGATCAGAG CTCCTGCAGA ACTGGGGGAGT TTACTTTTAC TATCTCTTCT
170581
170641
        CCAGGACAGG ACCTATCTCA AGAGACATGT TCAGAGTGAT TGCAACATAA AGAGTTTGCA
170701
        GACCCAAGGA GGTAGGGAAG GCAGAAAGAA GATGGGGGAG GCCAGGGATA GGCAACAGAG
        GAGTGACCAG GAGCGAAAAA GCCTGCCTCT TCTGAGAACC TAGCTGGGCT CTCCCTGTAC
170761
170821
        CCCCGATCCC TCCCCCCGC CCGCCCCCAC ACCCCTACTC CTGGGAGGTC CTCTAGGACA
170881
        GGGGCAGAGT CAGGAGGAAG TTTGAAGAGT GCCTAGAATA AAAAACAGTA ATTTAACTAC
170941
        AATTACCGGG TAGGCTGTTT TCCTCTCACA ATTTGATCAG TCTCTTGAAG CCACACAGAA
171001
        TTTCTTCTGA AGACGTGTAT TCCTTGGCAG GCTATTTCCT CCAGTGATAC ACCAGGCCCC
171061
        TCTCTGCTGG GGTCACTGCT CTTCTGGGGA GATGGGGCTC CCCTCCTTCC AAGGCTCCAG
171121
        GGTTCCTGTC CTGGGCCCCA CTCATCTAAG TTCTGAATCT TCTGAGATTT GGTGTAAAGT
        CTGGTGAAAG AAAGAGCAGG AAAGAGGTGA GAGCTGTAAA ACAAAGAAAG TCCTGACCAT
171181
        171241
        TACACACTCA CATATCCACT GAGAAAACCT TAGCCTGGAC CTTTTCCGTA ACCTTCACTG
171301
       CTCAGACACT TACATATTCG CTGCTAGTCC CCTCTGTTGC TGCCACTTCC TGGGTCAGGA
171361
171421
        AGTTAACTCA GACCGGATTA AACTGAGAAG TGAAACTACT GTGGGAGGCG GGGCTCATAA
171481
        GATTTAGGAG AAAACTAGTG ACGTTGTTCA TATCATTTGC ACTCCGCCTC TCCGGTAAAG
171541
        GAGGGGGAAA CGTAGGAAGA AAATATCCTT CTTTTACAGC AATAAAAAGA AGGAACCAAT
       TAATAACCCT GTAAACTATC ATGTGACCCC AACACAGAGT ATCTAAAAAC AGGAAGCCTG
171601
171661
        CAGAGGTTCA GTTCACAGAC TCTGATTTGA GATCTTTCTA CTTTTGCCAC CAACTCCCTT
```

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171721	GGGAGTCCTT	AAGCCTTCCT	AGCTGATGTT	ACTTCTTTTG	CTATTTATGG	GTTGCTTGTG
171781	GTTCTATAAC	TGCTCTGAAG	GGTGTGGTGG	AAAAAGGGGT	GGTAACAGCA	GTAGGACTCA
171841	TTGGCATCAC	AAAATTCATC	TGAGTCAGCT	TTCTATTCTT	CTCTGTCCCG	TTCTGTGTCT
171901	TGTTTTTCTC	CTTGCTGTCC	TTCTGCAGGA	CTCAGATCTT	CTTCAATAGC	GAGGGTCAGC
171961	CAGGATAGAA	AATGGGAGTC	ACTAGTGGCC	CAGCAGTGAG	TGCCCCCAGC	TTAGAGCTGT
172021	GTGGGATCCC	TGGGACCATC	ACTCTGCTTT	GTGCTTTGTG	GAGAAAAGGC	TGTGGGGTCC
172081	AGGGTCAAGT	CCTTAATGAC	TTAGCTCCAG	CTTCTCCACT	TCAAAATGAA	AGGAAAAGTA
172141	CTATCACCAC	CCGTTAGAAT	TATTATTTCA	TGGGGAAAAA	AGATGGATTA	CTATCTCACA
172201	ATAAGAGCTT	GTCACATTTA	TAAGTCTCAG	GTGTAAGAGG	CATTTATGAT	AACAACATAA
172261	TAAATGCTGG	CTTAAGTAGA	TGCAGTGGTC	CAAGGGAACC	AGTAAGGGGA	GCTCAGGACA
172321	CAGGTGGGAG	GAGAAATTAA	ACTTGAATTC	TGGGAGCCAC	TGGCCTGTCT	GGGCCCCTGG
172381	CCTGCCTGCT	GACCCTGATA	GCCAATGGAA	CATGGAGTTT	GGCCCAGCTG	CAATCCCTCT
172441	GGTCCAACTA	CTCAAAATAA	AGGCAAGATT	GGGAAACACG	TTCCTTTCTT	CCTATACCAA
172501	GCAGAAGACT	CTTCAGCACT	GCACCCTCCT	GGGTGCTCAC	AGAGCCTTCT	GTTGTTTTGC
172561	CACCTACGAT	TCATCATGCC	CTGGCATGAT	GGTTGCAGAC	CCCATGCATA	GCATGGGACA
172621	TTCTACTCCT	GAGGCAACCA	GCACACAGAG	AGAGGAGAAA	GAATGAGCCC	CTGAATCCTT
172681	GGTCCCACGA	TGAGTCCTTG	CAGATATCTA	CAACTTTCAT	TGTTGTGGAT	GTGACTCTGT
172741	ACCCAGGCAT	GGCTCATTCC	AGATCTGTCC	TATTGTCAGA	GGTGTTCAAA	CCAGAATGAC
172801	TCCATTTTGA	ATGGGGGCTA	GGTAAAATAA	GGCTGAGACC	TACTGGGCTG	CATTCCCAGG
172861	AAGTTAGGCA	TTGTAAGTCA	CAGGATGAAA	TAGGCAGTTG	GCACAAGACA	CAGGTCATAA
172921	AGATCTTGCT	GATAAAACAG	GTTGCAGTAA	AGAAGCTGAC	CAAAACCCAC	CAAAATCAAG
172981	ATGGCAACAA	GAGTGGCCTC	TAGTCATTCT	CATTGCTCAT	TATACACGAA	TTATAATGTG
173041	TTAGCAAGTT	AGAAGGCATT	CCCACCAGCT	CCATAGTGGT	TTATAAATAC	CATGGCGATG
173101	TCAGGAAGCT	ACCCTATATA	GTCTAAAAAG	GGGAGGAACG	CTTGGTTCTG	GGAATTGCCC
173161	ACATCTTTCC	CAGAAAACAT	ATGAATAATC	CACTCCTTGT	TTAGTACATA	ATCAAGAAAT
173221	AACTGTAAGT	ATCTGTATTA	GTCCATTTTC	ACACTGCTGA	TCCAGACATA	CCTGAGACTG
173281	AGTAATTTAT	ACCAGGAAAA	AATGTTTCAT	GCTCTTACAG	TCCCACGTGT	CTGGGGAGAC
173341	CTCACAACCA	CAGCAGAAGG	CAAGGAGGAG	CAAGTCAGGT	CTTACATGGA	TGGCAGCAGG
173411					CAGGTCTCAT	
173461	GACTATCATG	AGAACAGCAG	TATAAATTAC	TCAGGGAAAG	ACCTGCCCCC	ATGATTCAAT
173521	TACCTCCCAC	CAGGTCCCTC	CCACAATATG	TGGGAATTTA	AGATGAGAGT	TAGGTGGGGA
173581	CACAGCCAAA	CCATATCAGT	ATCCTTAGTC	CAGAAGCTGA	TGCTCTGCCT	GTAGAGTAGC
173641	CGTTCTTTTA	TTCCTTTACT	TTCTTGCTTT	CACTTTACTG	TGTAGACTTG	CCCCAAATTC
173701	TTTCTCACAC	GAGATCTAAG	AACCTTCTCT	TAGGGTCTGG	GTTGGGACCC	CCTTTCTGGT
173761	AACACTATCA	AAGGATCAGG	AAAAGGAAGC	TAGTGAATGC	TAAAAAGGAA	ACAAACTACC
173821	ATTACCAATA	ATAACAGCAA	GACAAAAGCA	AAACGGATTG	TGACAGCTGT	CCCATCTCAC
173881	ACCTGTTTCC	CATTGCAGGA	AGGAGGGGCT	GGTTCATGCA	CAGAGTGGCC	AATATTAGAA
173941					CAAAGGCAGG	
174001	AATCAACCTG	AACTATCCCC	AAGGAGGAAT	GCATTATCTC	TAATATGTAA	AGTTAGGCTT
174061	GATCCTGTGA	TTATGGGATA	TAGGAGTCCA	AAGACTCACA	ATGGGAAGTA	GGTCACTAGA
174121	GTCTCCTTCA	GAAGCTCTGT	ACTGTGTGTT	CCCACTGTGG	GCAAGAGTCA	GCACTCAGCT
174181	ATTCCTAGAA	TGCCTTTCCT	CAACTCCTTC	AGATTTTGCC	TCTCAACTAA	CCCTATCCTG
174241	ACCACTTGTT	AGCAAGTGTA	CCCCTCTCTC	CCTCCCAAAC	ATTTTCAAAT	CTATTTTGTT
174301	CCCATGGCAC	TTATCACTGA	ATATTTTACT	AATTTATTTT	GTTTAGTGTT	TGCTTCCCTC
174361	ATGAGAATGC	AAAGGGATGG	ATTTTTTCA	ATATTGTTCA	CTGATGAATC	CCAGTAACTA
174421	GAATATTTCT	AAGCATAGTG	ATGTGCATTA	AATCAAAGAG	TAACTTTCTG	AATTGCACTA
174481	AACACACATC	ACAAGAGGTG	TGTGCACATA	TGTGCATGAT	GCACGTAGTG	TGGTGTGGGT
174541	GTTGTGTGGG	GTATGTGGTA	CTGTGTGTGC	TGTGTGTGGT	ATGTGATACA	TAGTTTGTGT
174601	TAGTGTGATG	CATGTGATGT	GGTATGTGTG	TGCGTGTCCA	TACATATTAG	GGGTGGCGGG
174661	GATGTTAATA	TGTCAAATGG	TACTAGAAAG	TATCAGAACT	CATGGTGCTT	ACTGGTTTCC
174721	CAGAGAGCTG	CTTCTCTCCC	ACCTGTAGGA	TATACTGATG	GTTTGGACAG	AGAAGAAATA
174781	AAAAGAAGGC					
174841					CTTCTGGGGA	
174901	AAAAATCAAC	TCACAAATTT	ATTAACATGT	ACACAGGGAG	AACCATAGAA	TGATTATCCA

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174961				TATCCTGGCA		
175021	AGAAGAGAAA	CTCTGTTGAT	GGGATTACTG	TTGCGGATTT	TTGCTCCTTC	GCTCAGCTAG
175081				ATTAGGCATG		
175141				CTCAGCAAAG		
175201				AGGGCTTAAG		
175261				TAGACTGAGC		
175321	CCTGAACTGT	GCATGTGTTA	AGGAAAGGAA	TCATCCACTG	CAGGCATGTT	TAGGCAAGCC
175381				CATCCGGTGT		
175441	AGAGGTTCTC	TGGGTACCAT	TCCCTTACTG	TCTGCCTAAA	GCAAGCTGGC	CAACTCCTTT
175501	CATTACTAGG	GAGAGTAAGT	AGATCAGGGA	ACAGAGATTA	ACTTGAACAT	TATCTTGTGA
175561	AAGTCCGTTC	GGGCATGGTT	ACATTCTTGG	TCTTACAGGA	AGGGTAAATA	AAAATAATTG
175621	CTCTTTTTGG	TGGGTCTGGA	TCTTAGGTAG	ATAAAGAAAC	TTTAATTCCA	CGATGTGTTT
175681	TGGTAGGGAT	AGTTGGTGGC	AGGGATGTCA	GAGAGACTTT	GAGGCTTCTT	CAGTTCAATA
175741	TGACCAAGGG	CCATATATTA	GGGTATCAAT	TTCTGAGCCC	CAACAAGAGC	TTAGGAGAGA
175801	TGTGATAGCA	TCACAGTGTG	AAAGCAATTT	TTTGTTTGTT	TTTAGAGACA	GGCTCTTGCA
175861	CTGTCACCCT	GGCTGAAGTA	CAATGGTACG	ATCACAGCTC	ACTGTAATCT	TGAACTGGGT
175921				AGTGTTGGGA		
175981	ACCCAGCCTG	AAACTGCACC	CACTTTCTGA	TAAACTTTTC	AAATGACTAA	AGGGGAGAGA
176041	GTAAGCACTA	CTCAGAGGTA	GGAAGAAAGG	ACACAGGATT	ATAGGATTAA	AACAACAACC
176101	ACCAAAAAAA	ACCAGACCGG	TGTGGTGGCT	CACACCTGTA	ATCACAGCAC	TTGGGGAGGC
176161	TGAGGTGGGG	GGAGTCACTG	GAGGCCAGGA	GTTCGAGACG	AGCCTGGCCA	ACATAGCAAG
176221	ATGCTGTCTC	TATTAAAAAA	AAAAAATACC	TGCCTTGAGC	TAATCAGAAT	CATGGACCCT
176281	GACAAAGGAT	GTCCCAAAGT	AAGTCTTAGC	ATTTTTTTT	TTTTTTTGAG	ACAGTCTCGC
176341				GATCTCGGCT		
176401				TCCCAAGTAG		
176461				AATAGAGATG		
176521				TCTGCCCACC		
176581				AAAGTCTTAG		
176641				TTCACCCCAA		
176701				TCAACAGACA		
176761				AGAACAATTG		
176821				AATGTAACCA		
176881				ATTCCAAGGA		
176941				AGTTACATAT		
177001				AGGATCAAGC		
177061				TGGGATATTG		
177121				TATCTTATGA		
177181				TATAAAACAA		
177241				CAGGCTTCTT		
177301				AACATGCCTC		
177361				AACATTTTAC		
177421				ACTTTGGTCT		
177481				TTATACAAAC		
177541				CCCGCCTCTC		
177601				CCTCCCTAAA		
177661				ACCTCCTGAG		
177721				TCAAATGTTT		
177781				GCTCTGGAAG		
177841				CTAGTAATAA		
177901				TATGTCAACA		
177961				TAAATAAAA		
178021				TATAATAACT		
178081				TCCTGTTACC		
178141	CCAGACCCCA	AGAGAGGGTT	CTTGGATCTC	ACACAAGAAA	GAATTCGGGC	GAGTCTGTAL

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178201	AGTGAAAGCA	AGTTTATTAA	GAAAGTAGAG	GAATAAAAGA	ACGGCTACTC	CATAGGCAGA
178261	GCAGCTCTGA	GGGCTGCTGG	TCGCTCATTT	TTATGGTTAT	TTCTTGATTA	TGTGCTAAAC
178321	AAGGGGTGGA	TAATTCATGC	CTCCATTTTT	TAGACCATAT	AAAGTAACTT	CCTGACGTTG
178381	CCATGGCATT	CGTAAACTGT	CGTGGCGCTG	GTATGAGCAT	AGCAGTGAGG	ACGACCAGAG
178441	GTCACTCTCA	TCGCCATCTT	GGATTTGGTG	GGGAGCAGTG	AGGATGACCA	GAGGTCACTC
178501	TCATCGCCAT	CTTGGATTTG	GTGGGGTTTA	GCCAGCTTCT	TTACTTTTTT	CTTTTTTTTT
178561	TTTGCCCAGG	CTGGAGTGCA	GTGGCACGAT	CTCAGCTCAC	TGAAACCTCC	AATTTCTGAG
178621	TTCAAGCGAT	TCTCGTGCCT	CAGCCTCCCA	AGTAGCTGGG	ATTACAGGCA	TGTGCCACCA
178681	CACCCAGCTA	ATTTTTTATA	TTTTTAATAG	AGACCGGGTT	TCGCCATGTT	GCCTACGCTG
178741	ATCTCCAACT	CCTGCGCTCA	AGCCATCCAG	CCACCTTAGC	CTCCCAAAGT	GCTGGGCTTA
178801	TAGGTGTGAG	CCACCCCACC	TGGCCTAGCC	GGCTTCTTTA	CTGCAACCTG	TTTTATCAGC
178861	AAGGTCTTTA	TGACCTGTAT	TTTGTGCCCA	CTGCCTGCCT	CATCCTGTGG	CTTACAATGC
178921	CTAACTTACA	GGGAATGCAG	CCCAGCAGGA	CTCAGCCTTA	TTTCACCCAG	CTCCTATTCA
178981	AGATGGAGTC	TTTCTTGTTC	AAATACCTCT	GACAAGCCCA	ACACTTTGGG	AGGATGACAC
179041	AGGAGGATTG	CTTTAGCCTA	GGAGCTCAAG	ACCAGCCTGG	GCAACACAGT	GAGACCCCAT
179101	CTCTAAAAAA	AAAAATACAA	AAAAATTAGC	CAGGCATGAT	GGTGTGTGCC	TGTAGTCCCT
179161	GCTACTCAGG	AGGCTGAAGT	GGGAAGATGG	CTTCAGCCCA	GGAATTCAAG	GCTGCATTGT
179221	CAGAGGCATT	TGAACCAGAA	TGACTCTATC	TTGAATAGGC	GCTGGATAAA	ATAAGGCTGA
179281						GAGATAGGAA
179341		GGTACACATC				
179401	TGGCCAAAAC	CCATCAAAAC	CAACATGGCC	ACCAAAGGGA	CCTCTGGTTG	TCTTCACTGC
179461		TTAATTATAA				
179521		ATACTGCGGC				
179581	AACCCTCAAT	TTTGGGAATT	GTCCACCCCT	TTTTTGGAAT	GCTCATGAAT	AATCCACCCC
179641		CATAATCCAG				
179701		ACAGAGTAGC				
179761		ATGGACTTGC				
179821						AGGGACAATA
179881		TCTGAAGCCA				
179941		TCCCCGGGTA				
180001		TAAGACAGAG				
180061		GGTTTGAGAC				
180121		CTCAGTAAAG				
180181		CTGTTTGTAT				
180241		TGTACAGGAT				
180301		TGATAGGACT				
180361		CAGTGTTGCT				
180421		CTCACCACAG				
180481		CTTGAGACAA				
180541		AAAGATGAAA			- "	
180601		AAGTGGAGTG				
180661		ACGTTAATTT				
180721		TATTTGCCTG				
180781		ACCCAAGGGC				
180841		ACCTGGCATG				
180901		CTTAGTGAAA				
180961		TAATGGGAAC				
181021		GATATCAATT		**		
181081	GGATAGAACA					
181141	TCCGTTGCAA					
181201	TTCCTTCTCA					
181261	TTCTCTTGGT					
181321	TAAAAATTAT					
181381	GGGCAATGGA					
			TOTIGINGEL	-worker IWM	GOVITIGICA	TITININATO

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181441	GCGGCCAAGG	TTCAATCCTG	GCTTAGGGAA	TGAGTACTTT	CTGATTGATA	TCTGTGTGAC
181501		TGTTGATTCT				
181561		TGGGAGATTA				
181621						TGCTTTAGAG
181681		TATGGTTAAA				
181741	GACTCCTTGG	GAAAAGCAGA	GGAGGCACCA	CAGACCCCAT	TTTGGGAAAA	CCTCTGTTTT
181801	CCTCATGAAA	CCCCAGGAAC	TGGAAGTGGA	TAGATCCTTC	GCAAAATCTA	AGGCTCTGTT
181861	TGGCTTTGCA	TTATGTTATC	TGATGTTTTT	GACTTTTGGG	GGTATCAGAA	ATTACTTTGC
181921	ATTATGAGGG	AGATCTGGTG	TGTAATAACC	AGGTAGGAAA	TATACTTCTG	GGGATAGCTA
181981	AAGGCAAATA	TAGGTGAATA	CTTGGCTATT	TGCACTTTTG	GATCACAAGA	AGCATTCTCT
182041	TGACTACCTA	GAAGGTATGG	AAATGTCTCC	ATCCCCACCG	AGAGATAAGA	TTCCCAGGGG
182101	AGATGGCTGA	TCCCCCAAAA	GAGGGCTGAT	TCCCTCTTTT	GGGATCCAGG	ATCTGGTATA
182161	AAAATGGGAC	CCTGGCCAGG	CACAGTGGCT	CACGCCTGTA	ATCTCAACAC	TTTGGGAAGC
182221	CTCAGAGTTA	TGAATGTCTC	ACCATACTGA	CACTTTGTGA	CTGAGCTCCT	CTCTACCCTG
182281	GACACAAGAG	ACCCTAATAA	TTAGACAGGA	ATATCATTGC	CCCTATTTAG	TCTGAAGAAG
182341		CGGATCTTTA				
182401		AAATATGTCA				
182461		AAGGCTGAGG				
182521		GATAGAAGGT				
182581	GGTAACGGTA	AAGAAGCCAG	CTAAAGCCCA	CCAAAACCAA	CATGGCCACA	AAAGTGACCT
182641	CTTGTCATCC	TCACTGCTCA	TATACACTAA	TTATACTGCA	TTAGCATGCT	ACAAGACACT
182701		CCACGACAGT				
182761		GGGAAGAACC				
182821		CATTAGTTTA				
182881		TGCTGCTCTG				
182941		TCGCTCTGTC				
183001		TTCACCTCCC				
183061		GTGGGTGCCA				
183121		TTGGCCAGGC				
183181		GTGCTAGGAT				
183241		TTTTGTTTTT				
183301		GTGCGATCTT				
183361		TCTCCTGAGG				
183421		AGTAGAGACA				
183481	CCTCAAGTGG	TCTGCCTGCC	TCAGCCTCCC	AAAGTGCTGT	GATTACAGGC	ATAAATCACT
183541		CTTCTTTACT				
183601		TTCTTGTGTG				
183661		CTGGAGCAAG				
183721		AGTAAGAGGA				
183781	CTGTAAGAAA					
183841	AATCAGTGAA	CTCAAAGATA	GGTCAATTGA	AATGATCTAC	TCTGAAAAAC	AGAAAGAAGA
183901	CAGAATGAAG	AAAAAGAAAT	AGAGCCTTAG	AGACAGGGGA	TACCATCAAG	CATACTAATA
183961	TATGCATAAT	GGGACTCCTA	GAAGGAGAAA	AGTGAGAGGA	CAGGGAGAGA	GAATGTTTGG
184021	AGAAATAATT	TCTCAAAGCT	TCCCATGTTT	GGCAAAAAAG	CATTAACTTG	CATACATATT
184081	TTAGGAGCTC	AATGAATTCC	AAGTAGGATA	CACTCAAAGA	GATCCATACC	TAGACACATC
184141	ATAATCAGAT	TATCAAAAGA	TGAAGAAGAT	GAATCTTGAG	AGCAGAAAGA	AAGGAACAAT
184201	TCATCACATA	CAAATAGTAC	TCAAAAGATG	TCTGGAGTAG	GTATACTAAT	ATCAGACAAA
184261	ATAAACTTTA	AGATAAGCAT	TGTTATAATA	AATAAAGAAA	GGTATTTTGT	AATGATAAAA
184321	GTGTCAATTC	ATCAAGAAAA	CATAACATTA	TAAACATACA	TGCACCTAAC	AACAGAGCCC
184381	TAATATTCAT	GAAACAAAAC	TGACAGAATT	GAAGGGAGAA	ATAGAAAATT	CGACAATAAT
184441	AGTTGGAGAC	ATCAATACCT	CACTAGTTAG	ACAAGATCAA	САЛАЛАЛАТА	GAAGACTTAA
184501	CACTTGAAAA	CACCTAACCT	GACCCTAACA	TAAATCTATA	GGTCACTACA	CCCCAAAACA
184561	GCAGAATAAA	CATCCTTCTG	AAGCTCACAT	GAAACATTTT	TCAGGATAGA	CTGTATATTA
184621	CTTCATGAAA	TAAGTCTCAA	TAAATGTAAA	AGGACTATAA	TAATAGAGTA	TATATTCTCT

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						CEC > CCCCEC
184681					GGCGTGATGG	
184741					AGGTCAGGAG	
184801					TACAAAAATT	
184861					GGCAGGAGAA	
184921					GCATTCCAGC	
184981					TAGAAAAATA	
185041					CAGCCAAGAA	
185101					CATAGACACA	
185161					TACAATTCTG	
185221	AAAAACTCCC	CCCAAATGAG	GGTGAAATAA	GACAATTTAA	TACAGAGAAA	AGAGGAAGGA
185281					TACTGTATAT	
185341	TCTATTTCAT	TTAAAAAATC	AACCGTGCAA	TTAAATGGTA	GATTGTCTTG	CTTCTTTTTG
185401	ATTGACACAG	TCATTAACTA	AAATATTGTA	GTATTTTTT	ATCTCCCTGC	CTAAAGGCAA
185461	TAAACATCTA	ATCAGCAGAC	TAGAACAATA	TTTATAAAAA	TTTAAAAGTC	CTTTAGGCAG
185521	AATGATAAAA	GTCCCTTAGG	CATATTGAAA	TTCCTATTTA	TACAAAGGAA	TAAACAGTAC
185581	TAGAAATTGT	AACTATGTGA	GTAAACAGAT	AATATTTTTT	CTCCATAAAA	TGTGGTTGAC
185641	TATTTTCACA	AAAATAGTTA	ACAATGTAAT	GTGTGATTTA	TAGCATTTAA	AAGTAAAACA
185701	GGCCGGGCAC	AAAGGTTCGT	GCCTGTAATC	CCAGCACTTT	TGGAGGCCGA	GGCGTGCAGA
185761					GGCAAAACCC	
185821					TAATCCCAGC	
185881					TGCAGTGAGG	
185941					ACACACACAC	
186001					ATGAAAATAA	
186061					GATGTATACT	
186121					CAAGCCACAA	
186181					GAAAAAAAAG	
186241					GATGATAGAC	
186301					TCTAATACAA	
186361					GCAAAAAAAG	
186421					TAGTCTAGAA	
186481					TATTTATTTA	
186541					TGTTGCCCAG	
186601					GTTCAAGCGA	
186661					CATGCCCAGC	
186721					GGTCTCAAAC	
186781					ACCCAGCTCC	
186841					GATTTTGCCA	
186901					GGATGTTCTT	
186961					CAGTCAAGTA	
187021					TTACGCAGAG	
187081					GCAGCCTCAA CTACAGGTGT	
187141						
187201					TCACTTTGTT	
187261				· -	CTCCCAAAGT	
187321					TTAGCCTAAA	
187381					TCTCTCTCTC	
187441					AGAAGCAGAG	
187501					CTTTTATCAT	
187561					ATACCACCTT	
187621	and the second s				AATTTAAGTT	
187681					CTTTATTGTC	
187741						ATTAGAATGA
187801						GATTTCAAAT
187861	TTATTTCTAC	TGTAGTCAGA	TTTAATAATT	CATTTATTTT	TATTATTTTC	ATTTTTTTAG

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, , , , , , ,						
187921		TTTCTGTGTT				
187981						ACAACCAACA
188041		AAAGTGGGCA				
188101		ATATATGAAA				
188161		TTAACCTGAG				
188221		GGCTGGAGAG				
188281		GATGGCAGCA				
188341		TCTCGTGAGA				
188401		GTCACCTCCC				
188461		TGAGATTTGG				
188521		ATCAAAACCA				
188581		AAAAATAACA				
188641		AAATGTCAAT				
188701		GTGGTCACAG				
188761	AGAGAACAAA	TCTCTTGACA	TTACACAAAC	TGCATCTGGG	GCTAGTGGTT	AGAATATCCT
188821		GGTAGAAGAG				
188881	CACAAGCTGT	GTTCTCAGGT	TGACATATAC	TCATTTTAAT	AGTAAGAAAC	ACACCCTTGG
188941	GTAGAGAATT	AAAATGCTAA	TAATACATGT	GATGTATGTA	CTAGCGTGTA	TGGCAATATT
189001	GCATGCACAT	TCAAGAGACC	ACCCAAAACA	TATTTAACAA	CAATGCCCAT	TCCCACCCCC
189061	TCATGGATAA	TCACGTAGGA	CTCCCATAAC	GGGAGTTTCT	TCAGTGTCAA	TTGGTGCTGA
189121	AGTAGCCGAC	CCTGACTCTG	CTATCAGCGT	GTACTTTCAC	CTTGCAATAA	ACTCCTTTGC
189181		TTTGGACTGG				
189241		ACAACAGAGG				
189301	AAAATCTGCT	ACTGGCTATT	TATCCAAAGG	GAAGGAAATC	AGTATACAAA	GAGACACCTA
189361	CATCCCCATG	TTTATTGCGT	CACTCTTCAC	AAGAGCTGAT	ATATAGAGTC	AACCCTAAAT
189421	GTTCATTAAC	AGACAAATGG	ATAGAAAATG	TGGCATATAT	ACACAATGAA	ATACTATTTG
189481		GAATGCAATC				
189541		GATAAGCTAG				
189601		ATTTTTA				
189661		AGCAAGGAGG				
189721		AGAAATGAGT				
189781		GTATATTTTC				
189841		GTTTAAGGTG				
189901		TAAAAATATC				
189961		AAGAAAAAA				
190021		CATGCAGATA				
190081		GTTCAGATCT				
190141		AGGGTGTTAA				
190201		TTTTACTTTA				
190261		TTATGTATTT				
190321		TGAAATGACA				
190381		TTGCCCAGGC				
190441		TCAAGCGAGT				
190501		CCAAACTAAT				
190561		CGAACCTCTG				
190621		GAGACAGAGT				
190681		CAACCTCCGC				
190741		TACAGGCACA				
190801		CTATGTTGGC				
190861		GTAACACTCT				
190921		TTATGCTTTC				
190981		TTTTTATATT				
191041		GATTCTAACA				
191101		ATTTATTTTT				
		= - : -: <del>-</del>				

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SUBSTITUTE SHEET (RULE 26)

191161	GAACATAAAG	TGTGATAACT	GACATCCTTA	TTTCATTCCT	ACTCTGAGTG	GAAAGGGCAG
191221	GGGTGGAGAA	AGCATTCAAC	AATTTGCCAT	AATTATAATG	CTTTTTGTTA	CACTGTTTTC
191281					TAGGAGAAAA	
191341	TTTTCCTGGA	AAATGCCATA	ACCACGTCTC	TCAATTTTGT	TTCCATCTTT	CTTCCACATT
191401	TTACATAACC	TACATAAGAG	ACACATTATC	AAGTATATTT	TACATGGCTT	CTCAGTGTCT
191461	TCTCTGTCTG	CTAACAGGTT	TACCAAGAGA	TGGCACTCTT	GTATTTCTGG	TGGCTATGTC
191521	CATATCGTTT	TGCCTTTAAG	ACAGCGTAAC	TACTTCTTTC	ACCAGTATTA	AAGACATGTA
191581	CATTTGATCT	GGTTCTTGTG	GATGATTTTA	AATGACTCAA	GCTAATAATC	CTAATTTTAC
191641	CTAAACACTC	CATTATTTTA	AAATGTATTC	CTTTATGCCC	ACAATAAACA	TTTATTGACA
191701	TTAGGCTGGA	CATTAGGCTT	CTCTATGGCA	GACATTAGGC	TGGACCCTAG	CCATATATCT
191761	ATTGAGGGAA	TATTAAAAAA	TTTCTATATA	AGTTTCCAGA	AAGCCAAGAT	GTGTTTTAAA
191821					AAGAAAAGT	
191881					TGCTGCAGAG	
191941	GTGAACCATG	TTTTGGAGAA	GGAAAAGGTC	ACCAAGAGAA	GGAGGGGGTC	CAGGGTGTTC
192001	AGAAAGATTG	CATGCATAAA	GATCAAGGGT	AAAAAAAAA	ATTCCGTATT	ATGTAAATGT
192061	GAAGTTCCAG	GACCATGAGC	TTGGAGAGCA	TGAAGTACAG	GAGGAGGGTT	GGTTTCAAAT
192121	AAATCTGGGA	ATGAAACAGT	GAAGCCTCTG	GCAGAACTCA	CATCTCTTTC	CTCCCCTCTT
192181	CCTTGCACAT	TCCCTTTATG	GAGTAATTGC	AGGGATGGGA	AAAGTTCAAA	ACCACCACTG
192241	AGCCTAGGAA	GTGCTAGGGT	AAAGTGGAGA	ATGAACCTGC	GTGATTTGCT	CATCCTAAAC
192301	TAGGTTCTTC	TAGGAGAGCC	CTTCCCCATA	AAATCTGCCC	TCCTCGAAGG	GGCCCAGACA
192361	GCCTAAGCTC	ACCTCCCAAA	GACCCCTTAC	TTGCTGACTG	AATCTGATTC	CACCCAGACA
192421	TGGCCTAAAA	CCCTTCCATA	ACTCTATAGC	CAAATTCAAT	TTTAGACAGG	CCTCATACCA
192481	ACCTTTCTTC	CTCTAAGTCT	GCCACCCTAG	GCAATTCTCA	ACATTCTCTA	CACACTTTGG
192541	GGCCATAGAC	GTGCTACCAA	GTCTCCAGAC	CTAGACCTGA	TGGAGCAGTG	CTGTAATGAG
192601	ACGACCACTG	GCCTTTGAAC	CAGACCCTTC	TCTGTGGCTC	CTATGCATCT	CCAACCTGTT
192661	TTGAGCACTG	CTGCCAAGAC	ATCTTTGGCA	CTTTGTTGTG	AAGTTTTAAA	ACTGAACTAA
192721	TCTACAAAAC	ACCTAACCTT	TAAAAATTCA	TIGTCATITC	ATATCATGAA	AGATAAAGAA
192781	AGGCCAGGAA	ACTGTTCCAG	GTTAATAGAG	ACTAAAGAGA	TAGCAACCAA	ATGCAATTTG
192841	TGATCCTGGA	TTGAGGGGAA	AAAGTGTTGT	CAGAGACATG	ATTGGGACAG	CTGGTAAAAT
192901	TTGAATTTGA	ATTTAAAGAT	AAAGTATTGA	GTAATATAGG	AAGATGATTA	TCTGCAACTT
192961	TCAAATGTTT	CAGTAAGTAT	ATATATATAT	AAAGAGATAT	AAAGACATAT	AAATAAATGG
193021	ATAGGTAGAG	AAAAAGCAAA	TGTATAATAT	TAACAATCTA	GGTAAAAAGT	ATATGAGTGT
193081	TCTTTGTACT	GTTTTTCTGA	TTTTTCTATA	TGTTTGAAAT	CATTTTAAAA	TAAGAAGGTT
193141	TTTGGGTTTT	TTTTGTTTGT	TTTTTGTTTT	TAGAGACAGC	ATCTTATTCT	GTCACCAGGC
193201	TGTAGCTCAG	TGGCCCAATC	ATTGCTCACT	GCAGCCTCAA	CTTCCTGGGC	TCCAGTAATT
193261	CCCCCTACCT	CAGGCTCATG	AGTAGCTGGT	ACTTCAGGTG	TGCACCACTG	CACTCAGCTA
193321	ATTTTTATTT	TTTAAATTTT	TGTAGAGATG	GCATGTTGCT	ATGTCACCCA	GGCTAGTCTC
193381	AAACTCCTGC	CCCCAAGTGA	TCCTCCCACT	TTGGCCTCCC	AAAGTGCTAG	AATTATAGGC
193441	ATGAGCCACT	GCACCCAGCC	CCAAATAAAA	AAGTATTTTA	TTTTAATTAA	CTAATTAACT
193501	TTGAGTCAGA	GTTTCACCCT	TGTCACCCAG	GCTGGAGTGC	AATGGCATGA	TGTTGGCTCA
193561	CTGCAAACTC	TGCCTCCTGT	GTTTAAGCGA	. TTCTCTTGCC	TCAGACTCCT	GAGTAGCTGA
193621	GATTACAGGT	GCCTGCCACC	ATGCCCAGCT	· AATTTTTATA	TTTTTAGTAG	AGACGGGGTT
193681	TCAGCATGTT	GGTCAAGCTT	GTCTCAAACT	CCTGACCTCA	GGTGATCCAC	CCACCTCCGC
193741	CTCCGAAAGT	GTTGATGAGC	CACCACACCC	GGTCTAAAAA	. GTATTTTAAA	ACCACAGTCC
193801	CACTCTACCT	TGTCCTACAC	TACCAGGGGC	TAGGATCACC	CCATGTCITC	TAGGCTATGA
193861	GATAGAGGAA	TCCAAGGAAG	AAGATAAGCT	ACTTGGTTCC	TCTATAGGGT	CTTGTGTGTG
193921	CTCTCATGTG	CTCTCTCTCT	CTCTCTCTCI	CTCACACACA	CACACACACA	CACACACACA
193981	CACATGAATA	CCAGAGCTAT	CACTTTCCCA	GTCTAGTACI	CATCTCATCC	CAAGGGTTTT
194041	GTGTTGTAGT	GGTTTGCTCA	TTTCTTTGTI	TTGTTTGTTI	GCTTGGATTA	TTCTTTTTCT
194101	CTTTTTGCAG	CTGAAGGGAG	AATTTCCAGG	CCAGCCCTTT	GGCCATTAGA	GTTACAGTGC
194161	CTCTATTCAG	GCTTCATAGA	GAGACCTGGG	ATTCAGTAGT	GGGGGGCTTT	TATCCAGTTC
194221	AAAATAATGO	ATTCTCACCA	AGATGTACTT	TGAAATAAA	CAATACTAAA	ACACAAAATT
194281	TTATTTATGC	TGAACATTGA	ATCACTTTTT	TCTGTATTT	GTGTAGAAAG	TTATACACAC
194341	ACAAACACAT	TTGCTCCTGC	TTTGTTTAT	GGCCCAGGG	TATGTTTGGT	AATACTTCAT

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194401					CTAGACTCCT	
194461	TTCAGCATTC	TCCAGTGTAT	CTGTCATCTG	TCTACCTTAG	GATAGGGGTC	TCCAGAACTT
194521					ATATGAACTC	
194581					AATAAAGTAT	
194641					CTGAGTACCT	
194701					AGGGATAGGA	
194761					ACTACGTTCA	
194821					GGCGACAGAG	
194881					ATTTGAATGT	
194941					GCAGTTTTCA	
195001					CACATACCTG	
195061					GATTCTGAGG	
195121					CTCCTGCGTA	
195181	ACTGGGAAGC	CTACAGTTGA	AAATATTGGG	CTTGAGATCC	TGAAACAAAT	CTTGTATTTC
195241	ATTAAGACTA	ATATTTGGTA	CAGTGCAGCA	AATCAAGGGA	ATTTTGGTGG	CTGAGTTCTT
195301	TTAGAACTTT	TGCATTGAAA	TAGGTTCAAG	CAGCAATAAG	TTAAAACTAC	AACCTCAGCT
195361	AAAGGATTAA	AAGACACGTG	AGCTGGGTAG	GATGAGGTCT	AAGGTTGGGT	GTGGCGGCTC
195421	ATACCTGTAA	TCCCAGCACT	TTGGGAGACT	GAGGTGGGTG	GATCACTTGA	GGTCAGGAGT
195481	TCAAAACCAG	CCTGGCCAAC	ATGGTGAAAA	CCCATCTCTA	CTAAGAATAC	AAAAAATTA
195541	GCTGGGCGAG	GTGCCAGGCA	CCTGTAATCC	CAGCTACTGG	GGAGGCTGAG	GGAGGACAAT
195601					CGCACCACTG	
195661					ATAATAACAA	
195721					GAATAGTCTT	
195781					TTGAGATTAT	
195841					ATAGATGGAC	
195901					TTAGAGATGT	
195961					ATAGGCTCAC	
196021					CATGCCCCAC	
196081					TGTGTTATCT	
196141					CATGGATTCA	
196201					TTTCACATCT	
196261					GTGTACTAGT	
196321		CAAATTAGCA				TATTATTACC
196381					TTCCCTGCTT	CAAGCCTCAT
196441						ATTGTGGAAA
196501					GTTCTTTGCA	
196561					CAGTTCCTGT	
196621					CTCCCAGAAT	
196681						CCTGCCTAAA
196741					AGACAAGCTG	
196801	CCTGCGCTCA	AGCAATTCTC	CTGCCTTAGC	CTAAAAGTTC	TGGGATTATA	GGTATAAGCC
196861					CAGATGATTT	
196921	TCCATGGTAT	TCTAGGTTAT	TGTTGAGATG	GTCCTCTATT	GTCTTGTTCC	ATCTATTGAT
196981	TAGATAAAAC	GTTGTTCCTT	CTGTTATTTT	TCAACAGTAG	CTTTTATGTG	TCTCTCTTTA
197041	TCTTAAAATT	CTAACCAAAG	AGCTGCTCTT	TTCTTGGTGT	ACTTTACCTT	TGGTTGATCC
197101						ATGTGAGTCT
197161						TAGGTAGAAA
197221						TCTCAGCCAA
197281						GAACTCCCTT
197341						TCTTGGGAAT
197401						TGTGACCAGA
197461						TGCGAAGCTT
197521						TCCTGCTTGG
197581						GCACTTACTT

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197641		AGTGTCCTCA				
197701		TCATGTCCTA				
197761		GGGTTATCTT				
197821	GGCATATACA	TGGATTCAAG	TGTATGCCAC	GTGCATGTAT	TCCTTCATGT	ACTATTTCAT
197881	GTATTCTTTT	TCACATCTGT	TTTTTCCTCT	AAAATTTATT	TCCTTTTAAA	AATGAAAATT
197941		CTAAATTTGT				
198001	TTCCCGAAGT	TTTGAGTGAA	GTTAGTACTT	CAGAAAAACT	GTTTTGTATT	TTTCCTGTGA
198061		CTGCTGTGCA				
198121		CAAGATAAAG				
198181		ACTTCCCTAG				
198241	GCTTTGTGTT	TTCCTAAAAT	CAAAATAGGT	TTTTGCCTTT	TATGATTATA	CAGTAAATAA
198301		TGTGAAACTT				
198361	ATCTAAAAAT	CCTTGTGGCC	AGAATTAACT	ACCTTAGTTA	CTATTTTCTC	TATCTCTCTC
198421	TCTCAATGTA	TATTTGGTGT	AGGTATAGGG	GTGTGTGTAG	TGTGTGTGTA	TGTATATATC
198481	TGTTTCTATT	CCTGTATGTG	GATGTGCACA	ACGCATCCTG	CTTTGTACAC	TACAGTACTA
198541	GCATTTTTCT	AATGTAATTC	AATATTGTTG	AAAACATTTT	AAAAAAGCTT	GTATATATAC
198601		ACATACATGC				
198661		CACATGTATA				
198721		TTTTTTTGGT				
198781	GAGACGGAGT	TTTGTTGTCA	TTGCCCAGGC	TTAGTGCAGT	AGCGCGATCT	CACCTCACTG
198841	CAACCTCGAC	CTCCCGGGTT	CAAGCGGTTC	TCCTGCCTTA	GCCTCCTGAG	TAGCTGGTAC
198901		CGCCACCATG				
198961		GCCAAGCTGG				
199021		CTGGGATTAC				
199081		CTAAACTGTT				
199141		TATGTAAACA				
199201		GTCCCTTTAT				
199261		AAGTATTCAG				
199321		CAATGACTAT				
199381		GTTTGTCTTC				
199441		GGAAGAATCA			· -	
199501		AGTATAGTAG				
199561		TCTATCATTT				
199621		TAGTTTTCAT				TCTATTCCTG
199681		TCTACTATTG				
199741		ATTTCCTTTA				
199801		ACCAAGCCTC			_	
199861		CATACCACAC				
199921		CCCAGTTGGC				
199981		CATGTTTATC				
200041		CTGAACTCTT				
200101		TAATGTCATG			_	
200161		GCAGTGGATC				
200221		GGTCAGAAAT				
200281		AAAAAGTAGC				
200341		GGGAGGATCA				
200401		CTCCAGCCTG				
200461		TAATAATAAT				
200521		ATTATTATTT				
200581		AACGTGTGCC				
200641		ATGCATTAGC				
200701		CCCCAGTGAG				
200761	GCTCCCACTC					
200821	ATGTCAGGCC					
<del></del>						******

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200881					ATGTATACCT	
200941					GGAGTGAGGG	
201001	GCTACAGAGA	TCTTGATTGT	TGGTGGTGAA	GCAATGCAAG	AATTCATTCA	TTCAGTAAAC
201061	TAATGTTTAT	TAAGCGTGTA	CTGTCTTAGT	CTGTTCAGAC	TGCTGTAACA	AAATATCATA
201121					AGTTCTGGAG	
201181	TAAGATTAAG	GCCCTGGCAA	ATTTAGTGTC	TGGTGAGGAC	AGGTAGCCAT	CTTTTTGCTG
201241	AGTCCTAACA	TGGCAGAAGG	GTTGAATAAA	CTTCCTTGGG	TTTCTTTTAT	AAGGACACTA
201301	ATCCTAGTGA	TGAGGTTTCT	GCCCTCATGG	TATAACTACT	GCCCAAAGAC	CCCTCCTTCT
201361	AATATTATCA	CTTTGTGGGT	TAGGATTTCA	ACATGAGTTT	TGAGAGGATA	CAGACATTTG
201421	GATCATAGCA	CACACCATAG	GACAGACACT	GTGCCAAGAA	TTGTGGATAT	AGTGATTCTC
201481	AAAATGAACA	AGATCCCCTC	AGAGAGCTTG	CAAAATCCAG	CTATAAAATT	ATGCTTTTTA
201541	AACAAATTAT	GCAGTTTGAA	AAATCTACTC	TGAATCTTAC	TTGTGGCATT	GAATACTTTC
201601	GGCCACTCTT	TCCTTATTAT	ATTAAATATT	TACTCTTGTT	TGGGGGATCC	AGTCTCACCT
201661	ACTITITCIA	CCAGAACTGG	TATCAGCTCA	TGCTCTGCCT	TATGCAAATT	AAGAAAATAT
201721	CATACCTTTT	GGGTAAATTA	AGCCAAGAAA	GTTCTCCTTT	CTTCTCTTTC	TCTCTTTCTT
201781	TCTTTCTCTC	TTTCTCTTTC	TTTCTTTCTC	TCTCTCTCTT	TCTTTCTTTC	TTTCTTTCTT
201841	TCTTTCTTTC	TTTCTTTCTT	TTTCTTTCTG	ACAGGGTCTT	GCTCTATTGC	CTAGGCTGGA
201901	GTGCAGTGGT	GCAATCTCAG	CTCACTGCAG	CCTTGAACTC	CAGGGCTCAA	GCAATCCTCC
201961	TGAGTAGCTG	GGACTATAGG	CATGTGCCAC	AACATCAAGC	TAATTTTTGC	ATTTTTTTGT
202021	GGAGACGGGA	TCTCCCTATG	TTGCTAAGGC	TGGTCTTGGA	TTCCTGGGCT	TATGCGATTC
202081	TCCTGCCTCA	GCCTCCCAAA	GTCCTGGGAT	TACAGGCATG	AGCCACTGCC	CCTGGCCATT
202141	ATAACTATTT	TCATTGGCTT	ATCAGGCACA	TGATAACTAT	AATAAATCAA	TAACCAGAAT
202201	TTTTAAATAA	AGAAAGGAAG	GAATTGTTTC	AACTCTTCCT	GCTACCCCTC	TATCCCTCAA
202261	AAGGGTAGGC	TGAATGTTGT	CCTCCAAAGA	TATCCATGTC	CTAATCCCCA	GAACCTGTAA
202321	ATATATTACC	TTATATGACA	AAAGGGACTT	TACATGTTTA	ATAAGTTAAG	AATTTTGAGA
202381	TGGGCAGATT	TTCCTGAATT	TTGCAGATGG	GCCCTAGTGT	AATCACAAGG	GTCCTTATAA
202441	GAGACAGGCA	GAAGAGTCAG	AATAAGAGAA	AAATACTTCA	AGATGTTACA	CTGCTGGCTT
202501	TAAGGTGGAG	GAAAGGCCAA	GAGCCAAAAA	ATGCAGTGGT	CACTACAAGC	TGAAAAGAAA
202561	AAGAAATGGA	TTTTCCCCTA	AAGCCTCTGG	AGGGGGCACA	ACCTTGCCAA	TACCTTGATT
202621	TTGGCTCAGT	GAAACCCATT	TTGGACTTCT	GACCTTTAGA	ATTGTAAATA	TAATAAATAA
202681	TTTGTGTTGT	TTCAAGCCAT	CACAGTTGTG	GTAATTTACT	ACAACAGCAA	TAAAATAGAA
202741	TTAAATACAG	AGATCTGAGG	AGTTGAGTAG	GATAAGCCTA	CTCCAGCAGG	TTATTTCGGG
202801	AGTATGGTGA	GACTCACTAG	GATGGCGGAA	CTCAATTAAG	GAAGTCTGAA	GCTGATAAGC
202861	CAGAGAGGGA	AGGCTCTCAT	TTCATTTTAT	AAGGGTTGCG	TCACACTAGG	AAGATCCAAT
202921	AGCAACCACA	GTCTCAAAAT	TAATGATTAC	AAATAGGACA	CAATTCCAAG	AGTCGGGAGC
202981	CAAGCAGAAA	ATGGATTAGG	GAAGACATGG	ATGATATGAA	ACAGGAAGGA	GGGGTACAAG
203041	GCAGCTTCCT	GGGAAGTTGC	CAGGGCAGTC	ACAGTTCACA	TTCATTAGGC	TGTGGGCACC
203101	AAATGCATAT	GGAAAATCTA	GCTGACTTAA	CTGAACTCCT	GAAGAGGAAT	GAACACCTCA
203161	TTTATTGAGG	AGCTACTACC	AATTAGAATA	TGTATTTCAT	TTGTTCAATA	ACCCCATGAG
203221	TACAGTAACA	CAATCCTTGC	TTTACTAAAG	CGGAAGCCAA	TTCAAAGAGG	TTCAGTGACT
203281	TGTCCAAGCT	CAGGGAAAAC	ACTAGGAAGT	GAATATGGGT	CTGACTCCAT	CACTGATTTC
203341	AGGAGCCCTG	CCCTTTCCTC	CACACCATGC	CCCCTTGCTT	TCAGAAAAAA	AGGCTTGTTG
203401	ACTGAATGGT	TGTATGCACA	GTTCAAAGCA	GAAACACACG	ATGACATCTT	TTGAGATACT
203461	CTAACAGTGA	GAACTTGAAA	ATGAAGTTAA	AAATTAAGCG	GCAAAACCAA	GCCGAGGCTT
203521	TCTGAGAAAG	TGGGGCCAAA	CCTGTTGCCG	TCTGACTGCC	ACGTGGCTCA	CTATTTATCC
203581	CTGTAAAAAT	CTGCAAAAGT	ATTTGAAAGG	GAAGAAGGGA	CAGAAAACTC	CCTCCTTTTC
203641	CAAGTTAGCC	TTATAGTCTA	GGGCTTAAAA	TACTGGTTTA	ATGGTGAAGG	TAAGTGCTTT
203701	TCTTCTTTTT	GGGTAGAAGG	ATTATTACTA	ACTTACCAAA	GGTCCATTAA	GGGGAGGGAA
203761	CAGTTTTAGG	AGAAGTCAGA	GAAAAGACAT	TAACAGCAAC	ATAAGGATCT	CCATCTGGTA
203821	ATATTGCCTA	ATTCCAAAAT	GAAGAGACTC	TCTGAAAAAG	ATAACTGATT	CAATGAAGAC
203881	CCTAGGGCAA	GGCTTGAGAA	GCCACTGGTA	CCAATGGACA	CTGTGGACAA	TGGTCATTTC
203941	TCCAAGGACG	CTGTGAGTAT	TAACTGTGAT	GCTGTGATTA	GTCAGACTGG	GATTGGCTGT
204001	GGAATGAAAT	ACTGATCAGA	ACTGACAAGA	TTTGTGTTTG	GGACTGTGGC	TAACGAGTCT
204061	TTTCAGACTT	CTATATGAAT	TTGAAATGGT	CTCTCAGGAA	AAGGAGAACA	TGGCCGGGCC

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204121	TGGTGGCTCA	CGCCTGTAAT	CCCAGCACTT	TGGCAGGCTG	AGGCGGGCAG	ATCACTTGAG
204181	GTCAGGAGTT	TGAGACCAGC	CTGGCCAACA	TGGTGAAACC	CTGTCTCCAC	TAAAAATACA
204241				CTATGCGCAT		
204301				GAACCCAGGA		
204361				ACAGAGTAAG		
204421				AAGTTATGAA		
204481	TTGTACGCTT	GTAGAGATCA	CCTAGCTTGT	TGCCCTCATT	GTACAGCTAA	GAAAAGGCAC
204541				CAGAAAGATA		
204601				CTATGTTTTC		
204661				CTTTAAAAAA		
204721				TACATTTCAT		
204781				GTGACTGTTT		
204841				TTGCAATTTA		
204901				ACATTATCCC		
204961				TGGTTTACTC		
205021				AGGTTCATGC		
205081				AACAGAGATG		
205141				TTATAATAAA		
205201				GCGTATACCA		
205261				AGTAATTTTT		
205321				TACTTACAAC		
205381				GAAGGGTTCT		
205441				AGTGTCTCTA		
205501				TGTGCTTATT		
205561				GGTATGTTAG		
205621				GAAATTACCT		
205681				TTTTCATCCT		
205741				TTGTGATACA		
205801					-	
205861				TAATATTCAA		
205921				GATTAAAACT CCCATTGAGT		
205981				GTAGAACAAG		
206041						
206101				TTTCAGGCAA		
206161				GCCAGAGGAT		
206221				GGAAAGCTTT		
206221				CAGCCAAGCC		
206261				GGGTTTGACC		
206401				GATATCCAGA		
206461				TCCTGGTGCT		
206521				CAACAACAAC		
206581				ATACTATTCC		
206641				TGAAGTCCAA		
206701				CTACTAATTA		
206761	ATATAGAAAG					
<del>-</del>	ACTGGCACTT					
206821	TTTGTACACT					
206881	AAATAGCTTC					
	ATTTATCTCT					
207001	TTTTCATCTA					
207061 207121	TAAAGTACGG					
207121	AAAATAGAGG					
	TGTCCCCCAA					
207241	GTTCAAGCAA					
20130T	ACACCCGGCT	GITTITGTAT	TTTTAGTAGA	GATGGGGTTT	CACCATGTTG	GCCAGGATGA

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207361	TCTCGAACAC	CTGACCTCAA	GTGATCCACC	CACCTCAGTC	TCCCAAAGTG	CTGGGATTAC
207421	AGGTGTGAGC	CACTGCACCC	GGCCGATACA	TGTGTTTTTA	AAGTCACAGA	AATTTCAGAT
207481	GTCTTGAAGG	ATTTTAAGCA	ATTTAAAAAA	TAAAGTCATA	GAAGCTTCAA	TTTAGGAATG
207541	AATGGAAAAT	TGATGATATT	CTTAGGATAT	GGATTTTTCC	TAAAAGAAAC	AAATGTATGC
207601	ATCCCCAAAG	ATAATTTGAT	TAGTATACAA	ATATTAAATT	AAACATGTCC	ATATTTAGAG
207661	CCATGAATTC	TCTTTGCCTG	TCACAATAGC	TGGATTTATT	CACAATTGTA	GTAATTAGTC
207721	CCTGTTCATT	ATAATTTTCT	AGGTGATATG	AAGACTTTGT	CAGTCCAAGC	AAGTGTCCAC
207781	ATTGTGTGTA	GCAAACATGA	GAATAAACAT	TTTAAACTTT	TAAATGTAAT	ACATATTAGT
207841	GTTATGTAAT	GTCATCCTTC	ATGTTCGAAG	GCACATGGAA	CATTGTTCTG	GTGGTACAGA
207901	GGGGAGAGA	ACACCATCAG	AATGAAAGGA	AAGACCGCTC	TGGAACCTTC	CTCCTTAGCT
207961		GTTTAATTGT				
208021		TCTCTGTGGT				
208081		AAGGAGCCTT				
208141	TTAAATGACT	AGCTCTAGGT	CACACAGCTG	GAACTTACAG	CCAGATTTCC	TTTTAACAAT
208201	CCTGTAACCA	AAAGCATACC	AGTAGTGCCC	CATAAAATGT	AAGTTATAGA	GCTGTGTTGG
208261	GTCAAAACTT	TTACTGATGC	TAAGAGGAGG	CAACATTAAC	AAGGGGAAAT	TATTTGTGTA
208321	TTATGTTTTG	GATTATGTTC	TCTCCATAGA	TAAAAGACTG	TCGTAGTAAA	AGAGATTCAG
208381						AAATGGACGG
208441	GAAGCCTGCC	ACCAGGAAAG	GTAAAGCCAC	TGCTCTTGTT	TGCAGGCTAT	GTTAATAAGC
208501	TGAAGCTTAT	TCCGACACAT	TTACACATCT	CTGCATCACA	CTGACCCTTC	GTDARGATAC
208561		ACATTGGAGC				
208621		CTGTGAGAAA				
208681		TGGGAAAGTC				
208741		TGCAGTGGTG				
208801		GCCTCAGCCT				
208861	GATGAATTTT	TGTATTTTTA	GTAGAGATGG	AGTTTCGCCG	TGTTAGCCAG	CATCCTCTCC
208921		CTCGTGATCC				
208981		CCTGGCCCGG				
209041		ACCAAGCGGC				
209101		TTCCCAGACC				
209161		TATATTTTTA				
209221		AATAAGCAGG				
209281		GTTGTTCCAT				
209341		ATCTTCCAGC				
209401		TTTTTTTTT				
209461	GACATGATCT	TGGCTCATTG	CAACCTCTGC	CTCCTGGGTT	CARCECAGGEI	TCCTCTCTCTCT
209521		TAGCTGGGAT				
209581		TGGGGTTTCA				
209641	GATCCACCTG	CCTTAGCCTC	CCALANTGCT	GGGACTACAG	CCTCACCCA	CTCCACCCCA
209701	CGTAGTTTTT	TTTTTTTTT	AAGTTGAACA	TATGTGAAGG	CAGGACCTAG	TGACACACATAG
209761		TCCAAGTAGA				
209821	CCATCTCTCA	AATGTATTAA	AAGAGAATCC	TTGGATGTGC	AATACCTTAA	TTCNNAGCA
209881	GCTCGTTATG	TATAAACTCT	CAAGCTTTGT	CATABACAAA	TGTGCATAAC	ACATCCCA CT
209941	ATTCACTTAC	AGCCCAGGGA	ATTTTATTCA	CCCTCACAAC	CTTATCTCAC	TOCOTOTOCO
210001	ACTGTCATCC	CCATTCACTT	ראייייייייייייייייייייייייייייייייייייי	CARTATCACA	TARATGIGAC	ACATOTOCO
210061	TTTCTCTATT	TATCATGTGT	TTCCTATCCC	CTTCDAACAT	CCCCATATT	CCTTTTACTTC
210121	GTTATAAGAT	CCCATATTCG	CTCTCTTCAA	GCCAACCAAA	TARTTCACA	ANGTOCOTTO
210181		CTATITIGGT				
210241		AGCTGTGAGA				
210301	ATACATCTGT	GTTGTTGTCT	AGGTCCAGAT	TTCTCTTCAT	TACCOTTATCO	IGCICACIAC
210361	ATCATGCACT	TCTCAAACTT	CACCATGATA	ACCCACCCTC	TREGULATING	CATTGCGATC
210421	ATCGCCATGG	TGAACACCAC	TCAGCAGCAA	GGTCTATCTA	ATCCCTCCAC	TCALIGCGAIC
210481		CCTTCAATAA				
210541	ATGATGGAAA	ATAGGGCTCT	TICCAGCAIA	TOCATCANGG	CENTROLING .	CD TRACT TARGT
<b></b>			TIGITGWONG	AND THE STATE OF T	DURADURARU	CHIMONICII

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210601		AGTATGGAAG				
210661		GACATTGCTT				
210721	ATTATAAGGT	GGGGGAACTG	TAGAGTTAAA	TGTGAAAAAT	TTAAAAATGG	AACAGTTTAT
210781	GTGATGTCTT	CAATGAAAAA	CTAGGTATTA	CCTGGGCACA	TTCTTATAGG	TTACTCAATC
210841	CTATTCAGTT	CTCTGCCTGT	TTTATTGTTT	CTGAGCAATT	TTATATCCCT	GTAAATTCTA
210901		AGAAATGCAA				
210961	AGAAAAATCA	GTTAAAACTT	TTCTCCACTC	ACCTCCCAGT	TGAATTAGCC	AATTTTGCTG
211021	TTTGTTTGTT	TGTTTGTTTT	TTGAGATAGA	GTCTTCCTCT	GTCATTCAGG	CTGGAGTGCA
211081	GTGGCATGAT	CTCAGCTCAC	TGCAGCCTCC	GCCTCCCGGG	TTCAAGAGAT	TTTCCTGTCT
211141	CGGCCTCCCA	AGTAGCTGGG	AGTAAGGGGG	CATGCCACCG	CGGCTGGCTA	ATTTTTGTAT
211201		GACAGGGTTT				
211261	CCGCCTCGGC	CTCCCAAAGT	GTTGGGATTA	CAGGTGTGAG	CCACTGTGCC	AGGCTCTGCT
211321		AGTCTATTTC				
211381		ACCAGTTATA				
211441		TGTATGTTTA				
211501		TTTAAATTAT				
211561	ATTAATGATT	TTATTACATT	GGACCTAGCT	TATTTACAAT	GAGTACATAA	ATTTATTGTC
211621		CCTCCATTAT				
211681		ACCTTCAAGT				
211741		GCCTGCTTCT				
211801		GCAAGCCAGA				
211861		CCGATGAGAT				
211921		AGTTCAAGTA				
211981		CTCTGTTTTA				
212041		TTATTTATAG				
212101		CTTTATAGCA				
212161		AAAATCTATT				
212221		CAGGTAATAT				
212281		AATTATTTAT				
212341		GGAACTCAGT				
212401		GCACTGATTT				
212461		ATATAAGAGT				
212521		CAGATGAAAT				
212581		ATCATTTTCT				
212641 -		GCAAATTAGG				
212701		ATCAATATTT				
212761		TCATAATAGG				
212821		ATGGAGCCCA				
212881		TCTGATCCCA				
212941		TTTGCTGATC				
213001		TTTGGTCATC				
213061		TCTTGGTGGG				
213121		TTCAGGGAAT				
213181		AACGAAGCAA				
213241		TTGTCATCTG				
213301		GAGAGAAAA				
213361		AGATGCATCT				
213421		GGTTTCAGCT				
213481		CGGAAAAAGT				
213541		TACACCATCT				
213601		AGGAGAGAAA				
213661		TCTATTATGA				
213721	TTTTAAATTC	TTATTTTACA	GAGAAGGGAG	TTAAGGAAGG	TGGAGATTAA	GAAAATTGCC
213781	CAAATACAAA	TAGCCAGCAG	GTGGTAGGTC	TGAGATTTAA	GCCCATGCAG	ATTTTAGCCC

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213841	CAGAGCAGAC	ATTCTCAATC	ACTATGCTAG	ACTGCCTTTC	CATGGTATGT	GATCCTACTC
213901	AGGCCTCTAC	AGCTTTATCA	TTGCTGTTCT	CCCCAGCCTG	TCGTGCTGAG	AGTATATACT
213961	CGAAGAGCAG	AACTAAAATT	CCATCCAGCT	TCTCACTCCT	AGGTCCACTA	CACAGCTGCA
214021		TTTTACCTCA				
214081		CTCTATTTGC				
214141	TTCAACCTTC	TTCCCACCAA	AACCAAGTTA	GCTTGCTAAA	ATAAAGATGG	CACATTTTTA
214201		TGAGAATTTT				
214261		GAATACAAAA				
214321		GCATGGCTCC				
214381		CTCTGGCTCT				
214441		CCAAAATTCC				
214501		TTTTTTTTT				
214561	GACTGCAGTA	GCGCTATCTC	GGCTCACTGC	AAGCTCCGCC	TCCCGGGTTC	ACGCCATTTT
214621		CCTCCCGAGT				
214681		TTAGTAGAGA				
214741	TGATCCGCCC	GCCTCGGCCT	CCCAAAGTGC	TGGGATTACA	GGCGTGAGCC	ACCGCGCCCG
214801		CCTAAATCTT				
214861		TTATTATATT				
214921	CATTAGTGAG	TCATAAAATC	CATTGAGCGG	GTTAAAATCA	TTATTTTAAA	AAATGAATAG
214981	AATAGAATAG	AAATTGTTGG	AGTGCATTGG	ACATGGTAAA	GTTAAATATC	GATTCATGAA
215041	ACCATCGTTT	GAGGCATATG	TGTGTGGTTG	TATGTACAAG	TGTTTATGCA	TATTGGTGTG
215101	TGTGTTATGT	TACCCTGTAA	AATGCATTTC	TTACTATAGG	TCTCTGTGAA	ATATGTGTCT
215161		TAATGTAGAC				
215221	TTATTCACAT	TTTTCTCTCC	AATTGGACCA	GAAGCTCTTT	GAGGGCAGGG	GCTGTATCTT
215281	ACCGATTTTT	GTAAGTCTTT	CATTTCCTGC	CCCTAGCCTC	ATATTAGATC	ATGCAAGAAT
215341	GCAACTGTAA	TCACAAGAAA	ATGCTAATGG	GCTGTGATAG	CAGAGAGTTA	CTGTGACAAA
215401	CTAAGGGATT	TAGATTTGGT	CACATTGGTG	TTGAGGAGCC	ATTGAAGAAT	CAGAGAGTGT
215461		TTTGTTAATT				
215521	CTATTTTAGA	AATAAATACT	CTCATTGCCC	AATAATTCTA	AGTCTGCCAC	CTCACTGTTG
215581	GGACATTGTT	TAGGGAGGCC	ACGAAGTCTC	AGCCTTTGAT	ATTTTCATAA	GTGTTTTTCT
215641		TTTAGGGTCA				
215701		CTTGAGCTGG				
215761	AATCCTAACG	CCTCCATTTC	CTGAGCATCC	ATTTTGGCAC	CTACACCACC	CACATTCTTC
215821		GAAAATGTCC				
215881		ATCTAGTCAC				
215941		GTAGAGGAGT				
216001		TATCAAGCAC				
216061		AAGATGTCAA				
216121		ATTTCTGTCC				
216181		CAAATGTTCC				
216241	TGGCATTCTC	TCCTTTATGA	TATTTCCTCT	CTAGGTTATG	TTGGTGTGTA	ATTATTTATT
216301	TCTCCTTTTC	TTTCCACTAG	ACTGTGAAAT	GCTTGAGGCA	AGGAATCCAT	TCTATGTTTT
216361	CATCACTTGG	GTGTCATCAT	GGTGCCTGAT	TTTTAGCTTT	AAAATAAAAG	AATCAGTGAA
216421	TCCAGTAATT	AGAGGGGATT	TAAAGAAAAC	TAGTCCTCAG	AATCTTTTAA	CATAGAATGT
216481	TCTTCAAATA	AGGAATTCCA	ATAATAAGAC	AATTTTCTAC	ACTTGATTTT	GTTTTTATAG
216541		TCATTAAATA				
216601		TGTACATGTT				
216661		TGGCTTGAAT				
216721		TGATTTCATG				
216781		CGCAACAAGA				
216841		ATCATTGGAG				
216901		GAGGGCTCTG				
216961	TARARAGCTT	TAACAGGTTT	GTAGAAGGAT	TGAAAGAAAG	AATGGGAACA	TTTAGGTCCT
217021	TATGGTAGAA	TAAGCATTAA	TTGATTAGTG	TGTAGAAGGG	AGAGGCATGC	CACTTCAGAG

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217081						CTTCTTCCCA
217141						TGACCCCATG
217201						GGCTCAACAG
217261						GGTGTGGATC
217321					CTGATTAGAT	
217381	AGCCCAGTTC	TCCTGGACGA	GCTGTCCCCA	TAAAGGCGAT	GGTCACATGC	CTACCACTTT
217441					CACCATCATC	
217501						TTACTTCCTA
217561					TCTGTGTTAC	
217621					CCTTGAGAGA	
217681	TAATTCCCTT	<b>TTCACAGATG</b>	TGGAAACAGG	ACACTTAGAG	GTGAGATAAC	TTGCCCCAGG
217741	TTGCACAATA	CTAAGTGATA	GAGCTGCTGC	AGCATCCATA	TTCTTAACCA	CTATGCTATA
217801	CTACCACACC	AGCTGATTCC	AAAGCTTCTT	TTAGAAATAA	TATTGCTGGG	CCAGGCATGG
217861	TGGCTCATGC	CTGTAATTCC	AGCACTTTGG	GAGGCCGAGG	CAGGCAGATC	ATGAGGTCAG
217921					CATCTACTAA	
217981					TTCAGGAGGC	
218041	GAATCGCTTG	AACCCAGGAG	GTGGAGGTTG	CATTGAGCCA	AGATCATGCC	ACTGCACTCC
218101					AACCCAAGAA	
218161					TTATCTGAGA	
218221					ACAGTATCAT	
218281					GTATCTCTAT	
218341					GCCTTTTATT	
218401					GTCCAGGAAT	
218461					TAAGCGTGTG	
218521					ACAGCATGTC	
218581					TTTGTGGCCT	
218641					AACCTATGTG	
218701					TACCTGTTTT	
218761					GAAACTCTTT	
218821					CTGTCTGTGG	
218881					TTGGCCAGTA	
218941					GTATGCTGGG	
219001					ATTAGCTACC	
219061					CTCCTGTGCC	
219121					GCCCTATTGT	
219181					GTGACTATGG	
219241					TCTGTGTCCT	
219301					TGGTTCCCAG	
219361					GTTTGGCAAA	
219421					AATGACTTTG	
219481					GGCTTTCTAT	
219541	TTATAAAATC	ATCTTCCCAC	AGATATGCAA	GTTTCCTCAT	GGGAATCTCA	AGGGGATTTG
219601	GGCTCATCGC	AGGAATCATC	TCTTCCACTG	CCACTGGATT	CCTCATCAGT	CAGGTTGGGC
219661	CAGTTTATTG	AACATCTTCA	AGTGGCAGGT	ATTGTTTTAG	GTGTTGGAGA	TACACACGGT
219721	GCTCTAAAGA	TCTGGATGGC	AACACAATTA	CTCTATTTAC	ATGAGCCTCT	AAATCAGACT
219781	CTGGTAGGTC	AGATTTCCCA	GAGGAAGAAA	AATATAAGCT	TATTTTCTCA	AGATGAGAGA
219841					CAGCACGTGT	
219901	AGGTACATGA	GCATGAAACA	GTTCTTAGTT	ATGACCAGAA	TGAAAGACAC	ATGTCAAGGA
219961	ATAGCAAGAG	ACGAAGACAG	AGGGGCAAAA	GAAGATCATC	AAGAATATGT	TCAGACTAAT
220021	CCAATTTTTA	AAAAATCACA	AAAGGGAAAC	AAAGTGTCCT	AGGCCAGTTT	DDDCDTDDT
220081	TAATGTCTGG	AAACAGATCG	GCTGTGAGAC	ATTGCAAGGA	GGCTTGCTCG	CACAMAGGG P P
220141	ATGCAGGCTC	ATGAGGAAGA	TGAAAAGACA	GACCCAGGCA	GGGATGGAAG	GIGITIGGAA
220201	AACCAACTTA	CAAAGAGAAG	TTTTGTTTT	ACTACATTTC	TATGTGATCA	ACTTCCCACC
220261	TTAATATTTG	ACTAAACTGC	TAGGAATCCA	CTGTGACTAT	AATGCTGGAA	AGIICCCAGG
	· · · · <del>-</del>				GGAM	UTGWCIIWGI

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220321						ATTGAGATGG
220381						CAAACAGCAA
220441						CATTTTAGGC
220501		ATGAGTAATA				
220561		TGTTTAAAGA				
220621		CAATAGTTTC				
220681	TTCCTTGTGA	TCAACATTGC	AATACACAAC	TGGGAGGGCT	ACTAGAACTG	GTGTAGAAGG
220741	AACTTGTGAG	ATTGATCATT	TTCTCTGTTT	TTTACATCTA	GGATTTTGAG	TCTGGTTGGA
220801		TTTCCTGTCT				
220861	TTGGACAAGC	AGAACTTCAA	GACTGGGCCA	AAGAGAGGAC	CCTTACCCGC	CTCTGAGGAC
220921	ATAAAGTTAC	AAACTTAAAT	GTGGTACTGA	GCATGAACTT	TTTAAACATT	TTTTACTTCT
220981	CTCCATATTC	CTGACCATAG	ACTCAGCAGT	TCTTAACTCT	GGCTGTGTGT	TAGTCTTCCC
221041	TGGGGAGCCT	TTATAAGACA	CTGATACTTG	GGACCCACTC	CAGAGATTCT	GAATGAATTG
221101		GAACCCAGAT				
221161	CGCCCGGGGT	TGACAACAGC	TGGACAAACT	TGAAAAGTCA	ATTCATGTGG	CCTTTGAATT
221221	TTCCTCATTG	GAAAGTACTA	AATAAATAAA	AATTCATGTG	AAAATGATCA	CTGATAAATA
221281		GGGGCAGGTT				
221341		TCAGCACCGA				
221401		ATACATTGTA				
221461		TAAAATAAAAT				
221521		ATATTTATTT				
221581	CCAATACAAG	AAATAAA	TGAGGTTAAC	TTAGAAATCA	AGGACAGAGA	AGATAGAACT
221641		TATTGTGAGA				
221701		TATAGTTTAG				
221761		TGACAATTTA				
221821		AATGTATCAC				
221881		AGAAAAAGAT				
221941		CAGCGTCAAC				
222001		TGTCTCCTCA				
222061		ACCATATTTT				
222121	AATTCAGCAG	ATGAAGAGAA	TCTCCTAATG	CAAATCAATG	GGTATTTTTG	AGCAAGTTTT
222181		AGAGTGTCAG				
222241		TGACAACACA				
222301		GAAAGACATA				
222361		AAAAGGGTGG				
222421		CATAAAAGAG				
222481		CATTACAGAA				
222541		AAAATGAAAG				
222601		TACGTTACGA				
222661	CTAAATGTAC	AATTCCAGTG	ACTCAAACAT	TTTACTTAGA	AATGCACATA	TACATCCATA
222721		AACAATGTTC				
222781		AGAATGAGTA				
222841	GACCCCAATA	TATAATAGAT	GAATGGGTCT	CATAAGCACA	ATATTGATTA	AAGGAAGACA
222901	AAACGCACAT	TCTTTTAAAG	GTTTATAAAA	TACTTTTTAA	AAACAGCTAC	AACCAATCTG
222961	TCCTGTTAAA	AATCAGTGAG	CGATTTCCCT	TGTGCAGGGA	TGGGGGTTGT	GGCTGGATGG
223021	ATGGTACTTA	AGAAGTGCTC	CTGGGGTACT	AGAAATATTT	TATTTCTTGA	CTTGGATGTG
223081	TGTTTACTTT	GTGAATATTG	TACATTTATG	ATTTGTGCAC	GTTTATGAAT	GTAGAAAATA
223141	AAACAGAAAG	CAAATICAAA	GTATCATCCT	TTTGAGAGCT	TCTGCTCTGA	CTTCGTTTTG
223201	ACCAATGGAG	CAGTTGGGAA	GGGGTCTTGG	TCCTTCGGTC	CTTTGCTTTT	TTTTTTTTT
223261	TTTTTTTTT	TAGACAGAGT	CITACTCTGT	CGCCCGGGCT	GGAGTGCAGT	GGCTCGATCT
223321	TAGCTCACTG	AAAGCTTTGC	CTCCCGGGTT	CATGCCATTC	TCCTGCCTCA	GCCTCCCCAG
223381	TAGCTGGGAC	TACAGGCACC	TGCCACCATG	CCCGGCTAAT	TTTTTGTATT	TTTTAGTAGA
223441	GACGGGGTTT	CACCATGTTA	GCCAGGATGG	TCTCGATCTC	CTGACCTCGT	GATCCGCCCA
223501	CCTGAGCCTC	CCAAAGTGCT	GGGATTACAG	GTGTGAGCCA	CCGCGCCCGG	CCCCTGGTCC

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223561	TCTGCTTTCA	TGTTCTTCTT	GGTCCTGTTC	CTCCTCCTCT	TTTGTTGGAA	CTTCCAGTAT
223621	CAGAGCAGGA	. AGGAAGGCAA	TGGGTCAATC	GATGCTGTCA	GCTTTTGGAT	CAAACTGCAA
223681	GTTCTCAAAC	AGCAAAATTA	ATGAGCTCAG	GCTTTGAAGA	AACCATGACO	CTGAAAGCAT
223741	CAGTTGCTTC	CAATTGCATC	AGTTGCCACG	GGTGATAAGA	ACAATGATGA	CTCAGAATGC
223801	CTAGGTTTTC	CCAGCAGCTI	CTCTGAGGTT	TTCCCAGCAG	CTTCTCTGAT	TGATTCCTGA
223861	CAGATGACTT	CGGTGTGTCA	GACTTTCAGG	GTATCTTTCC	TTATGTGATG	GTTTGAGGAA
223921	GAGTTACCAT	TCACATTCCT	' AATGGCTTCA	GAATAGATGO	AATTGTGAAC	ТСАТАССАВА
223981	CATTTCTAAT	TCATCTCCCC	: TCCCCATCCC	TAAAGGATTG	TTTCTAACAA	TAGTCATGAA
224041	AATTAATTCA	CTTTTCTCAA	. ATAGTTTATT	GTCATCTACC	TAATGATGAG	ATGACTTACT
224101	TTTTCTCCTT	GACTGTTAAA	. TATTATGAAT	TATATTAATG	TATTTCTTAA	TGTTGAGCTT
224161	TCCCTTGAAT	ATTCTTTTGA	TGTACGACAG	AATTTGATTC	ACTAATAGTT	TATTTACCAC
224221	TTTGGCTGAT	GTACTGATAT	' ATGAGATTGG	CTCTGTATGC	ATACATGTGT	TTTGTGTATC
224281	TTTTTTGTGT	CTGGATATGG	AGCTTATGCT	GATTTCAAAA	ACAAGAAAGG	AGAACTTTCC
224341	TTTTTCCCCA	TTACTCTGAA	AAAGATTGAC	TAGAATGGAA	TTTTTTATAAT	ТССТСТТСТТ
224401	ATTTGAAAGC	TTGAAAGCAT	TGGTTTGTAA	AAATCATGCA	GGCTGAAAGC	CATTTTGAGG
224461	AGACTTTGAT	AACTTTCTCA	ATTTCCTTCA	GTTACTGGTC	TTTTAAGGGG	TTTTATATTT
224521	TTCTTTGATC	AATTTTGACC	ATTTATGTTA	TCTTGGAGGA	TCATCTATTT	TACACACTAT
224581	TTAAAGTATA	TTTGCAAAAA	TTCAACTGTT	TTATCAGGCT	ATCTTTTTAA	TAATATATTC
224641	ATTTTATCTA	TATCTGAGGT	TTTAGCTTCT	TTGTACTTCT	GACCCAATTG	CATGTGTGCT
224701	TTCTTTCTCC	TTCATTAGAC	TACTTAGTCA	TTTACTAATT	TTAAGAATAG	CTTGTCTTTT
224761	ATTTATTTAC	TTATTTATTT	TTGAGACGGA	GTCTCACTCT	GTCACCCAGG	CTGGAGTGCA
224821	GTGGCGCGAT	CTCGGCTCAC	TGCAACCTCC	GCCTCCCGGG	TTCAAGTGAT	TCTCCTGCCT
224881	CAGACTCCCG	AGTAGCTGGG	ATTACAGTCA	TGCACCACCA	TGTCTGGCTA	ATTTCTGTAT
224941	TTTTAATAGA	GATGGGGTTT	TGCTATGTTG	GCCAAGCTGG	TCTCAAACTC	CTGACCTTAG
225001	ATGATCTACC	CACCTTGGCC	TCCCAAAGTG	CTGGGATTAC	AGGCATGAGC	CACTGCGCCC
225061	AGCCCTGCTT	GTCTTTTTAT	TTTATATTTG	ATTAGCTTTA	TCTTTTTATCA	AGCTTATGTC
225121	CTATTTCCCT	TTGCTTTACT	TCATATAAAT	TTTGTTTTGG	ATAGTTTATT	TATTTTTCAT
225181	TTAATTATGA	AACAGGTTAA	AGCTTAGAGG	AAAATTGCTC	CTCTAAGTCC	AATTTTGTGG
225241	GCAGATTACA	TTTTGCTGTG	TTGTGCTCCC	AAATTCATTG	TTCTTTTAAT	GCTTTATTTC
225301	TCAAGTTAAT	AACCTATATA	GTAAAAAAGT	GGCTGTTGAC	TCTCAGCTTT	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT
225361	TTTTTTTTT	GTAGATACAG	GGATCTTGCT	GTGTTGCTCA	GGCTGGTCTG	AAACTGCTGG
225421	CTTCAAGGGA	TCCTCCTGCC	TTGGTCTCAC	AAAATGCTGG	GATGACAGAC	ATGAGACACC
225481	ATGCCTAGCC	ATGTCTCTCT	CCTTATATAT	AATAAGAAAA	CAGACACACT	GAGGCATCCT
225541	ATCATCTCAC	TCTTGGTTTC	ACTACTGTTC	TCTGGAAGTT	TTGCTCTGAC	CTTTTGCAGT
225601	TAATGTATTA	ATTTTGCATT	GAGTAGTTTC	CATAGAAGAA	TTATAGCATT	TGCATTCTGT
225661	TGGGTATTAT	ACTITICACT	GTTATTTGAA	CATAATTTGA	GGGCTGAAAC	CAAGATGAGG
225721	CAAGTGAGGT	GCCCAGGAAG	CAATATTTAA	GGAGGCATCC	TTTCTTAGGC	TCATGCAAGA
225781	ACAGAATTGG	CACATGAGAG	TGAGTGCCTC	CTTAATTTTG	AGTGCTGGAC	ACTTCTTGCT
225841	CACTTAGCAT	ACCCCTGGAC	AATGAAGTGT	TTTTTGTTTT	GTTTTTTCAT	GTCCATCCTT
225901 225961	TATCCTTCTT	CATCTCAAAA	CATTTCAATG	GAGTATTTTT	TTGGAGCAGT	ACTTGGATGA
	GCCTCTGAGT	CCCACAGTAG	CTGAGAATTT	ATTTCATAGT	ACTCTTTATG	ATCACTGTGG
226021 226081	AGCCTTAAAA	CATTGTAATA	TTAACTTAGC	TGGGAACAGA	AATTTTGTTC	CACAATTTGT
226141	CITATICAGA	ACAGTATTGA	CTTCCTGCTA	GTCTCTTCTG	ATGTCCAATA	TGAGGAAGTC
226201	TAGTTAGCCA	GCTACTTTTT	GTAGGAGAGC	TATGTTTAGG	CTAGGTGCTA	TAGGATTCTC
226261	TTTATCCTGG	AATTCCTTCA	CCAAGATGTG	CCAAGGTGTT	AATCATTTTC	TCTTGCTTTT
_	TGGCTGGTGG	TCTTAGAGTT	TCCTTCGATT	TTGTTTTATT	TAGTGATTGT	CCTCAATTTG
226321 226381	TTTTCTTTAC	TAAGAATCTC	TCTTCTATTT	ATCTGTATGG	TAAAACCTTG	TTGCCCATCT
226441	TTCTGGTTTC	TGCTGACTTT	CATTTTTGGA	CCTTTTACTT	TGCTTTCTCC	ATGGACTTTT
226501	TGGTAGTGGA	GGCAGGCAAA	CACTTTCCAA	AGTCTTTCTC	AATTTCCATC	AATTTCAACT
226561	TATTTCCTAA	AATTGCCTCA	GAATGTGCCT	ATGTCCACAA	TATCCCTCCT	TCCACTTTAG
226621	AAAGGAAAGG	CATUCACACT	TTATTTAGGT	GCAATGCCTG	AAGTGTAAAC	ACTTTCTGGT
226681	TTCACCTTCC	CTOTTC	CAAATATTGG	TTTGGGGATA	ACCTGCTAAT	GATTAACACA
226741	TAATCACTCA	CACTIGGTTT	GCCTGCTCCC	TCTTCTTTTA	TCTGCTGTGT	GTATTTTTT
/24	TAATCACTGA	GAATATGCAC	AGTATTGTAT	GTTTTATTAT	AAGAGAGGAC	TGGCCAGAGT

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226801	GGGAATGTT	C TGAATTCAG	ATABCTGAB	G CAGTACACO	3 M3 GG3 3 GG	A TTCTTTCAAA
226861	TGAAGCTGG	ATATTTTCC	AGAGCACCA	D CAGIACAGG	A TAGGAACTC	A AACTTGATAT
226921	GAATGATAC	A ATAAAGTGG	TAGAACTT	T ATTICAMIA	ATATTTAAA	A AACTTGATAT A TGAAATACTT
226981	ATTCTAATT	A TAGTCACTCT	ייי מייייטיטיטייי	T WILMMANIA	A ACTTATGTC	A TGAAATACTT
227041	TATTTACAA	ACAATTTATT	TTTTCATCA	1 ICAICITAT	A ACATGTTTAI	A TGTTTTCTTT A AAAATATTCA
227101	AAGGAATGC	TAAAGTTTT	LITTORION	R AAGIIITAG	A AATCAAGTT	AAAATATTCA A GAGTCTGAAG
227161	ACCATTTAG	TATCCABATT	. GTTTTNTTTTT	I IACAIGIIG	T ACAATCAAA	A GAGTCTGAAG A TTTACTATTT
227221	ATAATCCTT	AAAATTTGCC	TTACCACACAC	AAGCAGTAT	C CCTTCTAATA	A TTTACTATTT
227281	GCAGTGAGC	AACACAGTGC	CACTGCCCCT	AGAATTGCT	T GAACCCAGG	GACGGAGGTT GACTCTGTCT
227341	CAAAAAAAA	ασασασασά	. באכוטבטוו	CAGCCTCGG	C GACAGAGTGA	GACTCTGTCT CAAAAAAAATC
227401	CGCCTTAACZ	· ፓፓልጥጥሮጥጥር	TOTALARA	GCCAAAAAC	A AATAAACAAA	CAAAAAAATC CCCTTTCCTC
227461	TCAGCCCATT	GTCATATTT		1 ICTTTAAT	A CTACTAGTTI	CCCTTTCCTC GAGGTTTTTG
227521	TTTTTTTT	TTTTTGGAGA	TCCACTCTC	ACTIGCTIT	G TAGGACATAT	'GAGGTTTTTG
227581	CAATCTTGGC	TCACTGCAAC	CTCTCCCTCC	CICIGITGC	CGTGCTGGAG	TGCAATGGCG
227641	TCCAAGTAGC	TGGGATTACA	CCCACCCAC	IGGGTTCAA	CAATTCTCCT	GCCTCAGCCT
227701	TAGAGACGGG	GTTTCACCAT	GTTCCCCACT	ACCACGCCT	GCTAATTTTT	GCCTCAGCCT
227761	CACAATCCTT	GGCCTCCCAA	AGTGCTATCA	CIGGICICG/	ACTCCTGACC	TCAAGTGATC
227821	ATATATGTTC	ATTTTGAGTC	CTTTDACAA	TTACAAGCA	GAGCCACCTG	CCCAGCCAGA
227881	TCTTGAGAAA	ATCTCTGAAA	DGATCCCAAA	GTCATAAGA	TTTTAGGAAT	TCAGTTACTT
227941	TTTCATATTC	AGAATTGTTT	TTTNAAAAAA	AATTTGTAG	CAATTATATT	GATTTCTCTT
228001	TAAAGAAACC	ACCTGTGTGT	TCCTTARACC	TIGIATGIG	GAAGATTTTT	GCACTGTAGT
228061	GGGGTTACTC	TGAGAATCAA	JCC2222CC	ATAAGTACAT	GTATTCAAAT	AAATTGAGGT
228121	CTGTAGCAAC	TTGCTCCATT	CTTC A A A TA A	GAAGAAACAG	GCAGCCTCAA	AAGGTCTTAG
228181	CAACATTTAA	GGTCTCAGAA	CATARANA	ATAGGCTTGA	ACTTGTATTT	TCCCTCTACT
228241	TTTGCTTTAA	CAAACCCTAG	DEDECTOORS	TIGGIGAAAI	TTAAGTAAAG	TGCTCACTCT
228301	CAAAGGGAGT	TCAGGACACC	AGAGCIGGIA	GGCAGAGCCT	CAACAGACCG	TTTTAGCTTC
228361	GATAGTTCCA	CCADATADAG	TTCAAATCACG	ACCACAATAC	ATCACACATA GGTAAGAAAT	ATTGAGAAAA
228421	AGTTTATATA	AAATTTATTT	TTGAAATGCI	TATTCOTTA TO	GGTAAGAAAT GAATAGGACC	CTTGGAAATA
228481	AAGCCACCCA	TTTGCCAAAA	TAAAGTGAGA	ATCCTTOCTT	TTGGGGACTC	AGTTCTACTT
228541	CTCCAAGTGC	CACTAACAAT	TCTTAGGACC	TGACCTAGAS	GCCAGGTGAT	CTCTTTGTAG
228601	ATGATCAATT	ATTTCATTTA	AATGGCTCTA	TOAGCTATAA	GCCAGGTGAT	TTCAGTTAAT
228661	TCCCTGCAGG	GAACTGCAGT	GGCTTTTATC	AIGIGCAGAG	GGAACGGAGC	CCATCAGCAT
228721	AAATCACTTT	CAGGGTGGTC	ATGTAGTTGC	TTTTTTTTTT	TCAGAAGATG	AACTGTTTTG
228781	TTTTAATATG	TGACTCCTCA	GATTCAGAAA	GTGCTCCCTX	TCAGAAGATG GTCTTAAGAG	ATTCTGCCTC
228841	TCAGTGGTCC	AGCGCTTATG	AACCCACATC	Thaccomand	CCCTGGGGGA	TGAATTACCC
228901	AAATTGGTGC	CATGGACATA	AGAGGAAGGC	ACACTCARC	AGAGAGCCCC	ACTATCAGAG
228961	AATCAGTGGA	CAGCATCATT	ATTTACAACT	TTCTT ATCAC	CCAGGAGCAT	GCATGATGAA
229021	GCCAATCTGG	CACCATGAGC	TCTAATTT	CTTCCACTTC	TTGGAACCGA	GAAAATCCAG
229081	TGACTGTTTA	GCCATTTTAG	AGTGTGGCAT	ACCTCCCTCC	TIGGAACCGA	TTCTGATGAA
229141	TAAACGGGCC	TTTGCCCTCT	CTTATGAACA	TAGACACCAA	CTAAACTGTG	AGGTTGGATG
229201	TCCAAATGGT	GGCCTGAATA	CTATTTACAA	CTDACCAGGAA	ATGAAATTGA	TCACATAGGT
229261	TCCTCTTTTG	CAGATACCAT	CATTATTCAT	TIMOGIACA	AAAGTTAACT	GTAAGTCTTT
229321	GGTAATTTTT	AATAGAAATG	TAATAATTCC	TTCTCXXCTT	TAGTCTTTAG	ATTTGTATTT
229381	GATGCTCTCC ATGGGATTAT	ATGTCCTTCC	AAAAAAAGGT	ATCTCARGII	TAGTCTTTAG	TCTTAAGGTT
229441	ATGGGATTAT	TCCATTTTGT	TCTTTCTTAA	TATATACTII	CACCCACCAC	TCGCCTTCAG
229501	CTGGGTGAGA	TGCTATAGGT	ACAATGACAA	GTGATACCTC	GAGCCACTTT	TTTTGTGGCT
229561	TGGATAGCCT	AAGTGGTGAC	TTTTACCTCC	ACTCCAAATA	TATOTTOTT	GTCACAAAAG
229621		WCCWCICING (	GIGCTAGGGA	TACACCACTA	77777775 ·	<b></b>
229681	CCCUIGION	WWGWGWWIWW	<b>GACAATAAAT</b>	AAGTAAAGTG	Chromanam.	777 CCMCCCC
229741		WOMMMAN IN	AGCAGGCAAG	<b>ふににふつかつかつかっ</b>	CRESSESS.	~~~~~~
229801	. TELOCCCA1G	TATATATGTT (	CTATTGGTTT	TATTCTCTC	CACACCCCCCC	
229861	.u.cancilio	WGWWGTIWCI (	SGCTTTTTGAT	יייט איט איטידעיייי	A TOTAL COMPANIES	
229921	TCTINGIGIG	IALICAGIGI	TTTAAGAGAG	<b>プサイフ・ファック・</b>	330330333m	
229981	TTATCCAAAC '	TTAAGCCTTG	TTTAGGTAA	AAGGGCTCCT	CTTACARCACT	AGGACAAAAT
					C. INCAMBGT A	HGAAGGTTAT

# Figure 8 (Page 71 of 73)

220041	<b>Mamma</b>					
230041 230101	TATTTGGCA	T TTAAATCCA	A CTGAAGACT.	A ATAAGACTA	A TTAATTAAA	A GTTTTTAAAT
230161	CHCHMCIGG	G IGCAAAATA	A ATGGAACTG	C CATGCTCGC	ר אמפייפייפראי	T CACTCCTCTC
	ADADDDIAD	C AGCACGAAG	C TAATCCCAC'	T CATCTTGCA	G GTTGCTCCX	r rormanaans
230221	AMATCAGTA	A GACAGAAGC	T GGTCAGATT	A TCAAGAGCC	ר דבכדדם אמרי	CACCACMACC
230281	MILIIGGAAG	G GGTTGCTCT	C ATTAGGCAG'	T GCCTGACCA	~ `````````````````````````````````````	
230341	IGIAICIGA	A GCCATCATG	C CTAGTTATG	F TCCCCCACTO	יייי מייים מייים ב	CTCRRRCCC
230401	GGCCCCC1G	C ACCCTAGAA	A GCTGGGTGG(	F TTCTACTGT	ה שנות החווש השר	CTABABACCO
230461	ICIICIIIG	G ATCTGGACT.	I TACCTCTAT(	TGATTTTTT	ייי מידי מידי מידי די	TONTON
230521	CIGAGICIG	I CACTGCTGC	r aactcagca	TTCTAGGGT	TATTCCCCCX	TOOOTOLOL
230581	WWGWWIII.	C ATAGCTTCC	A GCATCCTCTC	TCCTTCATT	מיתי מבותיתיתים מידי ב	TONGO TONGO
230641	-A-11111C	I CIIGGIGIGI	GCAGCTCTC	CTCTCCTTCC	ייטיייטיתיטיתטיעטיי	TOOTHAMA
230701	CIMMCICCI	G CTTTTTTTCT	C TTTTTTTTTT	TTGAGACGG	CTCTCCTTCT	CTCACCACA
230761	CIGGWGIGG	A GIGGCACAAI	CTCGGCTCAC	TGCAACCTCC	GCCTCCCGG	TTCSACONS
230821	TOTOTIGOC.	r CAGCCTCCC#	AGTAGCTGGG	ACTACAGGG	CTCBCCBCTB	TOCOCCA CON
230881	AIIIIIGIA.	I TITTAGTATI	GCTGTCATCA	ATCCACATG	CCAGAAGCAC	CTRCRRS
230941	IMMITCITIC	- TAGGTATCAR	ACCCTAGGAC	TCTTTCCTCT	AATCACAATA	TATAATCCC
231001	GATICCCAA	A CACGGTCTT	TCATATACAT	TTTCCACTGT	י אראדארידירים מידער <sup>י</sup>	TCACCOCCA
231061	AGCICITACA	A CAAACACGCC	CTCCCCTAGG	AAGCCTTTAT	' אאאדנידידרירי	3CC33C33MC
231121	MOTOMOCOCAL	* CAGTGTCCTT	GTCACATCTT	' AGGTTCTACA	ביתידים עידיקיים	TTOTATIONA
231181	WIGINWICT (	CCAGAGGGTG	TTATCATCTT	TTTTTTTGAG	ATGGAATCTT	CCTTTCCTCC
231241	CCMGGCIGGA	I GIGCAGIGGC	ATGATCTCGG	CTCACAGCAA	CCTCCACCTC	CTCCCTTCS
231301	GIGHTICICC	TGCCTCAGCC	TCCTGAGTAG	CTGGGATTAC	AGACGTGTGT	CACCACACOM
231361	GGCIAATTTI	TGTATTTTA	GTAGAGACAG	GGTTTCACCG	TGTTGGCDAC	COMMICOROS
231421	AACTCCCAAA	CTCAGGTGAT	CCACCCGCCT	CAGCCTCCCA	AAGTGCTGGG	እጥጥ እ <b>ር</b> ለ ርርሞር
231481	IGAGCCACCA	TGTCCAGCCC	CATCTTTTTC	TTTTAGTTTA	GTTCTTAACA	ስ ስ ሞስ ርጥር <b>ጥ</b> ር እ
231541	CACAAAGIGG	ATATAACAAT	ATTTTGAATT	ATGAATAACT	AAATGAATAT	TTCCACATITE
231601	CCIGGIGCIC	TCAAAGTTTT	ATGTTACAAA	AGAAAAACAA	GTCTAAAATA	CCTCCCTCAA
231661	GITTIATCT	GTACTATGAT	TTCAAACCAA	ATAAAAAACA	GGTGGGGTAA	ABACTCABAC
231721	AGGAAATACA	TATAACTGAA	AAATTTTGGT	ATGTTAGTAT	GATAATACTA	<b>ここかい かかかかかか</b>
231781	CCIGITICCC	CAACTTCATT	TTCTATAGCA	ATAAAAAGAA	ACABGTAAAT	CTATATIONATO
231841	ITAATTTAAA	AGAAGTAGTC	TACCATCTCT	TCTGTTAAAA	AGAAAAAACT	ሽ <b>ጥጥጥጥ</b> እ እ እ እ
231901	ATTATCTCTG	GAAGGATACA	CAGGGAACAT	TGCTCTGGTT	TCTTCCDACA	CACAAAMCAC
231961	GAACIAGAGA	GCATGGCCAA	GTGGGGTTTT	GCTTTTGTTT	TICTUTETOR	A TOTOTOTO ACO
232021	TITITIATIAT	TITCTTTTGT	AGGTTTGAAT	TTCAAACCAC	ATABATCTCT	TACAMOOMOR
232081	INATAATAAG	TTTAAAATAA	AACTTTTGGC	TGGGTGCAAT	GACTTACACC	<b>ででする 3 でごごごろ</b>
232141	GCGC111GGG	AAGCAGAGGT	GGGAGGATAC	TTGAGGCCAG	GAATTTCACA	TCACCCTCCC
232201	CHACATAGIG	AGACCCTGCC	TCTGTAGAAA	TAAACAAAAA	TTAGCTGGAT	N TOOTTOOTOO
232261	AIGCIIGIAC	TCCTAGCTAC	TTGGGAGGTT	GAGGCAGGAG	GATCCTTTCA	CTCCACCACM
232321.	TIGMGGCIGG	AGTGAGCTAT	AATCACCCAC	TGCACTATAG	CATGGGCAAT	A A CCTCA CAA
232381	CITGICICAA	AAAAAAAAA	AGGGGGGGG	AAACAAATAA	ΑΤΆΔΑΤΆΤΑΑ	ACAAAA COOO
232441	IGITICAMAA	TATGTAATAT	TTAGCACTAA	AGAATTCTGA	ATTGTAGAGG	TABBARACORC
232501	TIMMAAGTTA	ATAATTATTG	TCTCCTTTAA	$\Delta \Delta C \Delta \Delta T T C T T$	3 TO 3 3 3 CO 3 D	• • • • • • • • • • • • • • • • • • • •
232561	CHONWANTCH	TCCATATCAG	CAAGCTAAAC	TTTCTCAAAA	TGACATATCC	A TOTAL A TOTAL O
232621	CICCCAGGIA	MITAGCAGGC	AGCCTCTACT	CAGGTTGAGT	<u>እጥጥር ርጥ</u> እጥር	T
232681	MAN TO CHOUSE	IGCICCAAAA	TCGGCAACTT	TTTGAATGCT	<b>ልልሮልፕሮልምም</b> ሮ	TORRACOROM
232741	GCICAIGGAA	IATITCAGAT	TTTGGATTTT	TGGDTTTGDG	カサカベサベスペサス	M3 3 M00 0 3 3 3 0
232801	WI I CCWWWIC	TGAAAAAATC	TGAAATACTT	CTGGTTCTAA	GCATAAGCCA	TROTORROOM
232861	GIGILAGCIA	ATTAGACCCT	TCATGGTCTC	TTCTAGACCT	CDCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	7 7 CCT3 7 CCT
232921	CIMICCICAC	TTCTAATAGC	ATGAACTTTT	CTGTTTTAGA	ATAATTTCCA	<b>ででででとるととる</b>
232981	TO TACKNOW	AIAGIACAAA	GACAGTACAG	GAGAGTTCCC	יייייייטייטייטייעייטייטייעייטייטייטייטיי	C X C C TT X C C TT TT
233041	TCCCCCATTG	TTAGGATTTT	ACATTATTAT	GATACATTTC	תמשמשמשת מ	CCAACTCACA
233101	TIGHTWCWIG	AAACTCTATT	AACCAAACCC	TAGACTTTAT	CTCCATTTCA	CCXCTCTTTC
233161	CACIMAIGII	TICITICIGT	TCCAAGGTCC	AATCTGGAAT	ACCA CACTCC	y manaramana
233221	CATATCTCCC	TAGTCTTTTT	TTGTCTGTGA	CAATGTCTCA	GTCTTTTCTT	GCTTTTCATG
					<del>-</del>	

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233281	ACCTTAACAG	TCCTGAAGAT	CATTTGCTTT	TTTTTCATAA	TTACACCGGA	GTTATAGATT
233341	TTTTGAAATA	ATACCACAAG	GGCAAAGGGC	CCTTCTTGTC	ACATCATTTT	AGGGAGAACA
233401	TGATATCCAC	ATGACATCAC	TGATATTAAC	CTTCATCATG	TGGTTTAGGT	AATGTTTCAG
233461	GTTTCTCTAC	TGCAAAGTGA	TTTTTTTCCC	TTAATTTAGC	CCACCTGAAC	TTATCAATTT
233521	TGTTTTCTTC	CATGACTAAT	ACTTTTGTTA	TTATAGCTAA	AACTTCATTG	GGGCCAAATC
233581	TTAGATCATG	TAAATTTTCT	TCTATATTTT	ATTCTAAAAG	CTTGTAATGT	TTGATACATT
233641	CTAAAAGATG	TAATGTTTGA	TACATTACAT	CTAGTCCTTT	GATTTATTTT	TAGTTACTTT
233701	TGTATAAGGT	GTGAGAGATG	TCTCCAGTTT	CACTTTATTA	ACACATTGTG	GTGTTCCAGT
233761	ACTATTTGTT	GCTAAGACTA	TCTTTTTTCC		TTTGCCTTAG	TTGGCAATAT
233821	TTTTGTTGGT	TTATTTCTAG	ACTGTTTATC	TCATTCCACT	GATTTGTGTC	TATCTTTTTG
233881	ACAAAACTGT	TGATTACAGT	AAGCTTTGAA	ATAGTTCATT	TTTTGTGTCA	ACTTGACTGA
233941	GTCAGGGGAT	AACCAGCTAT	CTGGTTAAAC	ATTATTTCTG	GCTGTGTTTG	TGAGCGTGTT
234001	TCTGGATGAG	ATTAGCCTTT	GAATAGGTGA	TCCTAGTAAA	GTAAACTGTC	TTTCCCAGTG
234061	TGGATGGCAT	TATGCCACCT	GATATTCAGG	GTCTGAATAG	AAGAAAAGGC	AGAGGAAGGG
234121	GGAATTTGGG	CCTTTTTTTC	TGCCTCACTG	CTTGAGCTGG	GACATCTCAT	CTGGTCTCCT
234181	GCTCTTGAAC	TGGGATTTAC	ATCATCAGTT	CCTCTGGTTC	TCAGGCCTTC	AGATTCAGAC
234241		CACCAGCTTT		CAGCTTGCAG	ATTACAGATC	ATGGGACTCC
234301	TCATCTTCCA	TAAATGCATG	AGCCAATTCA	GTCTATGTCC	TTGAAAACTG	CCCCACTGCA
234361	GATTAAGGCT	TTTTTCCACT	AGGTGAAATA	AAGAAGCTTG	TTAGACAGAT	TTCCCTTCAT
234421	CCAGTGCCCT	CTCCTCTTTA	AGTTACAACA	CATTGGCTAC	ACCTAAGTGC	AGGGGTGGGG
234481		AGTCCTCTTG		AGAGAACTGT	ATTGGGAAAG	CTCTAGAAGT
234541	GTTTGATACA	TACATAAACA	AGGCATGGTT	TTTGCACTTA	ATTTCACATT	ACATTTTTCC
234601	CAGAAAAAAA	GGAATGTATA	GGCATCACGT	AACTGTACTA	GCTGGAGTCA	TTCTTCCTGA
234661	TTATCAAAGG	TAAACAGTTA	TTAATCCTAT	ACCAAGATGT	CAAGGAGAAG	TACTTTTGGA
234721		TTCTCTGGGA				GTTAATGAAA
234781	AACTATAGTA	CCTTCCTATA	AGCTGGATGA	CTAATTACCA	GGCTCATTTA	GGAATTTGCC
234841	TTACCAAGTA		GCAGCTGAGG		AGACAAATGG	AGCATAGAAT
234901	AAGAGTAGTA	AAGAATGCCA	AAAATGCTGT	CATGTATCCA	TTGACAAAAG	GAGCTATAAA
234961	GCCTTTAGGT	ATTTTCACAC	TTGCTCTGTT	ACGTAAATGT	ATGTGTGTGT	GTGTGTGTGT
235021	GTGTGTGTGT	GTG				

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1	CACACACACA	A CACACACAC	CACACACACA	CACAAATGA	G GTATATAAA	GGTCTCCTAA
61	AAIGICATCI	GATATTTGT	. ATTTCATATT	CTCAGATTT'	TAATCCATT	r Accraccror
121	ATTITAGATA	GCCTTGTCTC	AAACAGAGCI	'GGGACCTGA'	r gagtgaaaa	r GAGCTCACCA
181	GAAGAAAAA	CAAACAGGC	A TTTCAGAGAT	TGAGGCCAAG	S AAGTTAAATO	מאת התדידים ב
241	GCAGAGCTTA	A GCTGCTTGAT	: GTGAAAAGAG	ACCAGCGTG	G CTGGAACAGO	DAAGGAGAAC
301	AGCAGAAGAG	GTGAACAGAG	GCCAGAGATO	GTCACTGAG'	ממקררר מידים ז	CTCNTCCTN N
361	GGAGTATGGA	. GAATGAATTA	. TTGCATGTAT	'TGAATATGT	GGTGACGTG	CTCACACATA
421	CTTTGGATT	' GTAGAGATGA	. Aggaaatgta	GCAAGTGAC	A CTCTTAGAA1	CTTCATTCA
481	GTAAATGGTA	GTGTCAGTTA	. TTGAACTGGG	GAGAACTGG	AGGGATAACA	CCCTTAACCA
541	GCACGTTTAT	TCCTGTGTCT	' TGGAAGTGTT	'TAGGGTGAAI	GACCTATTAC	ACTO CTARRO
601	GGAGATGTCA	AGTGAAAATG	TGGCTACACA	CATTTGCATT	TCAGAAAAA	CCTCNCCCTC
661	GAGATGTAAA	ATTGGAAGTT	'TACTGCATAT	AGATAGTCTT	TGGAACCGT	GTATTGATCA
721	AGCCATTAAT	GAGACAGAAC	AAAGACTAGG	GACCAGAGCC	AAGCTCCAAG	ייי או או אוייי איייייייייייייייייייייי
781	TTAGAGGATA	GTATAGTCTG	GTCATTTTGA	GGTGAATACT	TAATAACAGA	ስር እ አጥጥሮ <b>ረ</b> ጥ
841	GAAGTGTAAA	TTTAGAGCCC	TACACTTTTA	GCTCTGACTA	TTABCCBATA	CACCAAACAA
901	TGGATATGGT	TATCTGCCTG	GTGTCTGTGA	AATAATTTAA	GCCAGGAAGA	GATCCTCACC
961	AGAAACTGAC	TATGCTGGCA	ACTIGGATCT	TAGATTTCCA	GCCTGCAGAA	<b>ででごででかご カカ</b>
1021	ATAAATGTCT	ATCGTTTAAG	CCACCAGTCT	GTAGTATTTT	GTTATEGEAG	TOCARCOTOR
1081	CIAAGITITIG	GTACCCAGGC	GTGGGATGCT	GCAACAACAA	ATACCTABAC	ATGGGGAAGT
1141	GGCTTTGGAA	ATTGGTGATG	GGTAAAGGCT	GGAAGAGTTT	GAGGTTCATA	CTAGAAAAAC
1201	CCAATTGTGA	AGGGACTATT	GAAAGAAATA	TGGACATTAA	AGGCAATTCT	GGCDDDGGCC
1261	CAGAAAGGAA	GAGAGCTGGA	CAGAAAGCTT	CCATTTTCAT	AGAAACTTAG	ስምምስ ምስ አ <i>ር</i> ር
1321	ATCATGGATA	GAATATTAAA	TATGCTGGTT	AAAATATGGA	CTTTAGGCCA	GGCGTGGTGG
1381	CICACGCCTG	TAATCTCAGC	ACTTTGGGAG	GCTGAGGGCA	CAGATCACGA	GGTCGGGAGT
1441	TTGAGACCAG	CCTGGCCAAT	ATGGCGAAAC	CCTGTCTCTA	CTAAAAATAC	DADATTACC
1501	TGGGCATGGT	GATGTGCTTC	TGTGGTCCCA	GCTACTCGGG	AGGCTGAGGC	TGAAGAATCC
1561	CTTAAACCCCG	GGGGGTGGAG	GTTGCAGTGA	CCCAAGATCA	CACCACTGCA	CTCCAGCCTG
1621	GGATACAGAG	CAGGACTCCA	CTCCCCCCGC	CACACACACA	CAAAAAATAT	ATATATATCC
1681	ACATTAAAGT	CAACTCTTGT	GAGGTCTCAG	ATGAAAATGA	GGGACAGGTT	ATTEGAAACT
1741	GIAGAAATCA	CIGITCITGT	TACAATGTGT	CAAGAACTTG	GCTGAATTAC	CCTCTACTCT
1801	TTACTGGAAA	GAACTTATAA	GCAGTAAAAC	<b>TGGATATTTA</b>	CCAGAAGAGA	ТСТСТАВССВ
1861	AAGTATTGAA	GGTGTGATTT	AGGTCCTCCT	TACTGCTTAA	AGTGAAATGT	GAGAGGAAAG
1921	AGCCGAAATA	AAGAAGGAAT	TTTTAAGCAA	AACACAATCA	GAACTTGGAG	ልጥጥርርርርአጥለ
1981	GATTTCTCAA	TCTATATTGT	AAAAATTGAG	AAAGTTTTTC	TTGAAGAGGT	ATGGTTGAAC
2041	AATGTTTCT	TITTCTTTTT	TTTTCTTGGT	TTTATTTTTA	The	מיי איי איי איי איי איי איי איי
2101	GGGTCTGGCT	ATGTCATCCA	GGCTGGAGTG	CAGTGGCACA	ATCTCAGTTC	ACTCCAACCT
2161	TIGCCTTCAG	GCTCAAGCAA	TCCTCCCACC	TCAGCCTCCT	AAGTAGCTCC	CACTACATOR
2221	AIGCACCACC	ACACCCTGGC	TAATTTTTTG	TTGTTGTTTA	TAGAGATGGG	CTTTTC A CAT
2281 2341	GIIGCCIAGG	CTGGTCTCTA	ACTCCTGAGC	TCAAGTGATC	TGCCCTCCTC	ACTOTOCONA
2401	AGIGITGGGA	TTACAGGCGT	GAAACACTGA	GCCTAGCCTG	AACAACCATT	TCATABACAC
2461	ATAMIGGGIG	TGACCCAAGG	ATTTAATCAG	CCATCTCAGC	ACAACCCACC	77C7C7C7
2521	GGATTATTCC	AGCAGAGACA	CTGCCAATTT	AAACTAACGT	AGGCAGAGAA	AACBCAAACC
2521 2581	AMCAMAGGAA	GGTTGTCGAC	TTTTTGAATT	CTATAGAACA	GGATCATAGA	COTTA COTTOCO
2641	TOTCHAIGIG	TACTATTCTT	TAAGAAAAGG	AAAGACTGAC	CCACCAAAGG	מאס מייייים מאס
2701	GATCACTAGG	GCTGACTCTT	TTTTGTTTT	TCTTGAGGCA	GTCTCACTCT	CACCCACCCT
2761	GINGGGCAAT	GGTGTGATCT	CAGCTCACTG	CAATCTCCAC	CTCCCAGGTT	CAAGGGATTC
2821	TCTTGCCTTA	GACTCCCAAG	TAGCTGGGAT	TACAGGCTCT	AAATCTGTAC	CCTCCCGAGT
2821 2881	AGCGCTCCTG	CCACCACTTG	CCCAGCTAAT	TTTTGTATTT	TTAGTAGAGA	TGGGGTTTTCA
2941	CIMIGITGGC	CAGGCTAGTT	TGGAACTCCT	GACCTCCAGT	GATCCATTCT	CATTGGCCTC
3001	CCAMAGIGCT	GGGATTACAG	GCAGGAGCCG	CCAGGGCTGC	CACTTTGATG	TCAGACTCAG
3061	AGAGIACAGA	TGGGATAGGG	TGGGGGTGGG .	AACATGTAGT	CAAGGCTGAC	サンス へいかいかか
3121	I CAAAGATGC	CCTGCAGAAC	TGTGTGGGAG '	TCTCTCACAG	ATGGCTGCCT	GGGTGGGACC
3181	CONCOMME	GAAAGACCGA	GACTTCAGGC .	AGGGCAGATG	GAGTAGGCCA	<b>カウザカクカクカクク</b>
	CAGAGGTGAC	HCTGAGACAC	CACTGGGCCT	GGAAATCAGG	GCATCAAGCC .	AAAGAGGGTT

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3241	TO THE CONTRACT OF THE COMMENTAGE
3301	
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5761	GGCATTTCAA AGTAGAAGGT AAAGTATTTT GGAGATGAGG AGACAGGACA GAGCTACGAG GAATGTCCTT TGCTCAGGGA CTAGCCTCTTT AGACAGGACA GAGCTACGAG
5821	
5881	
5941	
6001	
6061	
6121	
6181	
6241	
6301	
6361	
6421	GCCAGGCTGG TCTCAAACTC CTGACCTCAG GTGATCTACC CACCTCAGCC TCCCCAAGTG
	TOTAL SIGNICIACC CACCTCAGCC TCCCCAAGTG

Figure 9 (Page 2 of 74)

6481	CTGGGATTAC	AGATGTGAG	A CACCAGATC	A GCCTCAGAA	3 ACATTTTCT2	TTGGAAAGAG
6541		**************************************	A TIAGICIAA	ירות אידיידים ו	T T A ROTH COMMA	
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7261		CITIMAGGIL	3 LITETTAGT"	ייייטיע ערוייטיטיר	3 3 3 3 mmana	
7321		TINGULIGI.	LUGALIAAATT	אידי אידי את אידי בעם		
7381		WOINGIGC !	L AATAAACCAT	יוייטי עידייתייטיתיטיתיט	CRCCCCC	
7441		TIGGIGAGG	AGGITCTTT	ייטטעעטייירעט אינייי	TOROGRAMA	A3 = 2 A4 A
7501		- TUCCAVCAV	7 ALTAAGCCCT	ACCECTARGE	7000001100	
7561		~+********	LITCCAAGGCC	САТАВСССВВ	TOTAL	
7621		TWANDOWN	Late I delicated at the care	ここひ ひり カヤククク		
7681	ATGTGAAACC	AGACCTTCAA	ATCTGATGAT	TCTCACCCCA	GCTGCCCATT	CCCAAGAGCC
7741	TAATTTAAAA	ATACCCTCGG	AAAATTCTAA	TATCTCCCCA	TCAAAGGTGA	AGAATCGTTG
7801		TAGILLICAL	CCAAATGGGA	ר אדררא א רבר		
7861	-1100012200	AGGGGG I GGA	GGGAGGGAAG	AGCCCANNNC	CCMCCAMOGG	
7921	GTGTCAGTAT	CTGGGAAGTG	GGAGGCGCGT	CACCACTAAA	CAGCTTCTGC	CCCCGAGCCG
7981	ATCTCCTGCC	ACACACTCGG	ATTTGAAGGC	TCCAAACCAA	ACAATGCAAA	TAGGATTATT
8041	GGAGTTCCAG	AAGCGTTAGA	CTABACGACT	GCCTCTCTTT	GGCCAGTCTG	ACGCTTCAGT
8101	CGCAGATGCA	TAGGCAAGAC	TTAGCCCGCC	TAGACTUTTUTC	TGCCCACTTA	AGCAGCTGGG
8161	AAGCAGAAAC	CGGCCGGGCG	CGGTGGCTCA	CCCCCCTCTTAT	CCCAGCACTT	ATTCCGATCA
8221	AGGCTGGCGG CCGTTTCTAC	ATCACCTGAG	GTCAGGAGTT	CGACACCACC	CCCAGCACTT	TGGTAGGCAG
8281	CCGTTTCTAC GTCGTCTGAA	TGGTGGCGGG	CGCTTGTAAT	CCCATCTACT	CCGGCTAACC	TGGTGAAACT
8341	GTCGTCTGAA	CCCGGGAGGC	GGAGTTTGTA	TECRETCACC	AGGGAGGCTG	AGGCCGGAGA
8401	CCAGCTTGGG TGCAGAAACC	CAACAGGAGC	AAAACTCCGT	TTCDDDDDDD	CGAGATCGCG	CCACTGCATT
8461		MODULLCOR	AGAAAACCTC	GGCGAGATTC	XCXCXXMOG-	
8521	TCTCTAGAAA C	TTTGTCCATG	GTCCCAGATC	TCCATTTCTT	ACAGAATCCA CECCO	GGAAAATAGG
8581	CAGATCCCTA (	GAAGCAAAGG	TTTTTTGGG	GGACCCTCTC	GIGGGIGGG	CAGCTGTTAC
8641	GGGCAGTGGC	ACGATCTCGG	CTTACTACAA	CCTCCCCCC	TCACTGTTGC	CCAGGCTGGA
8701	TGCGTCAGCT	TCAAGAGTAG	CTGGGAGTAC	AACCEARCE	CCAGGCTCAA	GCGACTCTCC
8761	TTTTATTTAT	TATTTTTATT	TAGTAGAGAG	CTCTTTCACC	A TOTAL CONTROL OF	CAACTTATTT
8821		~~ 1 CAGG 1 GA	IL AGGUT TOTAL	CCCCCCCCCC		
8881			THURSTICK	1 C1 CCCCCCC	^	
8941	GTAATGGCAA (	CCTAGACGCT	TGAGCTTCTT	ACAGGCCCA	CTCTATTTCC	TTTTCTGCCT
9001	CCGTTCTACA T	TAGGGACAT	TAGTCTGTTT	TRERENCE .	AGTAAGTTGC )	ATGTCAGGCA
9061	TTTAGGTAAT A	TACTCTGCA	CTTTAGCAGG	AATCCAACACC	TTTCAACTCC (	CTGGTTAACT
9121			CIAAATTTTTTTT	י הפיחור או את את ממ		
9181						
9241			CHACKE ATTOR	א מותיידיתיזבוו	7M/7MM2	
9301		~~~~~~	CIAGAATATT	ו שור איר איושושויין ו		
9361	TACTATTACA T	ACAAACAGA	CCAACCTTTTA A	TTABCACC -	ATGATAGTAC C	TAACTGACC
9421						
9481	TACCTACAAG C	AGTGAGGTT	AGCTCTTCCT T	TENACTIAC (	TRAGTGCCT T	TTTTCCTTC
9541		WING COLLL	CCGGGGA(G("""" Z	י שאשאא מידעאו	18.3 2 mas	
9601			CITAGG AGA Z	CCDCCCCCC C	'^	
9661	CCCTGAGCAA T	GGTCACCCG	GCCTAGCAGT 1	TETTERESCE	GATGTTAGG A	AGGACGCCG
			INGCAGI	TGTTGAGCT (	CICGICGIT G	CGGATGGCC

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9721	AGCTGCAA	GT GGCGCGGG	AT GATGCGAGT	C THE CHARLES	T 0000100	C GTTGCCGGCC
9781	AGCTCCAG	GA TCTCGGCGG	T CAGATACTC	T ARCACCAC	r CGCGAGCCG	C GTTGCCGGCC C CGGCGCGCCT
9841	GCCCCAAC	CC GCTCTGCGT	A GTTGCCTTT	AACACCGCC	CCAGGTACA	C CGGCGCGCCT G GCCCACCGGG
9901	<b>AACTGGAG</b>	AC CAGCGCGAG	A AGAGCGGGA	T TTCCCTTTTC	GGTGCACTC	G GCCCACCGGG F GCCTCCTTGC
9961	TTACCACG	TC CAGACATTO	C AATCAGACA	1 11CGCTTTG	CGCGAGCTT	C CTAAGCTCAC
10021	GAGAAAACA	A ACAAAATCA	A GAAATATCT	A AAAICACCA	AACCAGCAG	C CTAAGCTCAC G GTAGTTCCTG
10081	GGGAGTAA	T CCGACTTT	T GATTGGTCG	A AAACATGGC	GCTTTTATA	GTAGTTCCTG  GCCAATAGAA
10141	AAGCTGTAC	T TTCATACCT	יב פאנוטטונטיי	G CTCTCCCC	TAGTCAGAT	A GCCAATAGAA TGTGTAGTTT
10201	GTCTTCCA	TAACTAAGA	G GTACTCTCC	A WCCCMCA	GGATGACAA	TGTGTAGTTT CCTATAAGTA
10261	GCAGAAATC	C GCTCTTTAC	T TTCGbCbCb	r TUCUTUATTY	GCATAAAAG	CCTATAAGTA CTGAGCCAGCC
10321	AAGTCTGCT	C CCGCCCCGA	A GARGGGGGG	TICIGGIGI	TTAAGATGC	TGAGCCAGCC GCAGAAGAAA
10381	GATGGCAAG	A AGCGCAAGC	G CAGCCCCAA	AAGAAGGCAG	TGACCAAAGO	GCAGAAGAAA GTACAAGGTG
10441	CTGAAACAG	G TCCATCCC	D CAGCCGCAA	GAGAGTTACT	CTGTGTACGT	' GTACAAGGTG
10501	TTCGTTAAC	G ACATATTTG	A CACIGGCAIC	COCCAAGO	CCATGGGCAT	GTACAAGGTG CATGAATTCT
10561	AAGCGCTCG	A CCATCACCT	C CAGGGAGAGA	GGCGAGGCT	CCCGCCTGGC	CATGAATTCT GCATTACAAC
10621	GAGCTGGCC	A AGCACGCCG	T GTCGGAGAT(	CAGACGGCCG	TGCGCCTGCT	GCATTACAAC GCTTCCCGGA
10681	AAGTAAACA	T TCCAAGTAA	CCTCTTARCY	. ACCAAGGCCG	TCACCAAGTA	CACCAGCTCC
10741	CCAGATACC	C ACTABABGA	CTGTCCCCA	CCTAACCCCA	AAGGCTCTTT	CACCAGCTCC TAAGAGCCAC
10801	ATTAGAATG	T AGGAACTGG	A GACCCCTCCC	ACGCCAAATT	TTATTTGGCG	GCGGAGGGGT
10861	GGGTCCTGA	A CCCGAAGA	a GCCACCCAmm	GACAAGTGTT	GCAGCTTAGA	GCGGAGGGT GAGGGACAAA
10921	TTAAATTTA	A AATGGGGAC	y ACCURCATI	AAAAATGGGT	TTGGGGTCAA	TTCGTTGTGC
10981	CAGGCTCGC	T TAGGTTTCA	AGCGGCCATT	TTGCTAACTC	GGCGTTCCCG	TTCGTTGTGC GAAGAAACCG
11041	GTTGCCGTA	A TGTCATAATT	ACCCAGCTGT	CTGTCCCTGT	CTACGTCGCC	AGGATCAACG
11101	AAATATTAA	C CAATCGAGG	TCGCCACCAG	CTTCTAGCCA	ATAGGCTGTC	CTGTCATTTT
11161	TGGGAACCT	GGCAGTAAC1	AAAGCTGTTT	TGAGACTCTG	ATTTACATAG	CGGACCGGAG
11221	CTTCGTAGT	A TACTGAAGG	GCCTAAGGAA	GGACTCCCCC	TCTGTTTTCG	TGGCGCACAC
11281	TAACCTAATA	A TGCGTCAGTT	TGTGTCTCCT	GGGTTTCCAA	CTGCCCCGGT	AATAGTCTTT
11341	GCACTGCGC	AGATGTTGCT	TTGATAACAA	CACTAAGGCA	GTACAGAACT	Aaagatgtaa
11401	AAATCAAATC	AAATTTTGCT	TCATACATCT	TATTCTATTC	AACTGGTTTA	TTCAAGATTC
11461	TGATTGAAAC	TTAAAATCTC	TGAATCCCAG	TGCTCAGTCA	GCCATAAATG	GTGTGTTGCC
11521	TTTAGGAGAZ	GCCAACTCTT	CGTAGGGGGC	TIGIAACATG	CAGAAAAGTT	TGAAAGTTGC
11581	AAGGCCAGTA	AGGACTAGGC	AACTGCTGGG	TAAATTGACA	AGCCTTCGAA	CACTGAACTG
11641	AGCACATACA	CTGTGTCTCC	GCTGGGTGGG	GGAGAATGAA	GAGGAGACGT	CATTAAACTT
11701	TGGCCTGGGA	AATTCCACAT	TAGAGGACTC	TCCCTTCCTA	GACAACTGCA	GGCCGCTTTG
11761	TTTTAAGATG	AAGGGTTAGA	TCCCTTAAGT	ATTTTACTCA	TGGTCTTTTC	CAGGTAAAGA
11821	TTTAGCACCT	AGAAGTTTGC	CGTAGTCTAC	CTATCTTTT	ATTCAAGTCT	AGAACACGTT
11881	AGTGTTAAAG	CAGATTTTTA	TTTCTCCATT	AAAAACCGGG	AATATACAAT	TTAAAATTA
11941	CATAAGGACT	GTGTGATCTT	CAAACTTAAA AAATCTGCAA	TACCATGTAA	TTTAGGTTAC	AGTTACTTAA
12001	CCTGTCTTTG	GTGCCAGACA	AGGCCTTATA	CTTCTTTCAC	ACCTGGGAAA	TAAACTAAGG
12061	CTTGCCTAGA	TAACTTATCT	GAGAAATTCT	CITGAACACT	GCTGTGCAAT	CACAGGCTGC
12121	CAAGTAAATT	TTTTTTTCTT	TTTTTTTTT	GATGAGAAAT	GAAATTTCCA	GAGTCCCTCA
12181	GGCTGGAGTG	CAATGGCGCG	ATCTTGGCTC	ACROCAROCT	AGTITCTCTC	TTGTTTCCCA
12241	ATTCTCCTGC	CTCAGCCTCC	GGAGTAGCTG	CCATTAGCACCT	CCGCCTCCCG	GGTTCAAGCC
12301	CTAATTTTGT	ATTTTTAGTA	GAGACGAGGT	TTCTCCATTC	CATGCGCCAC	GACACCCTGG
12361	TCCGGACATC	AGGTGATCTG	CCCGCCTTGG	CCTCCCATGT	CGGTCAGGCT	GGTCTCGAAC
12421	CCACCGCGCC	GGGCCTAAAT	GGTTTTTTT	TTTTCTAMAG	TCCTGGATTA	CAGGCTTGAG
12481		- SOME SOME	GUILITTAN	י מחשים אוארים אירוים וויים		<b></b>
12541	TCTGCAAACA	AAATTCAGTA	TTCTTTCCCC	CCCTTTTTCCC	ACTAATCAGT (	STAACCAAAA
12601			AAATAATTU	י <del>רושהייטירייטירייטיטירייטיריי</del> סירידיר ב	mana	
12661		UTTWILL	TUTTGGTCAT	ب لا حاصب ساساس لا بلسا	Million Casaa .	
12721		AUTUCAUTIVE.	GAACCAACAG	ABBTCTCTCT /	70111111111111111111111111111111111111	
12781			CIGITUACTO	TOTOTOTO A	mmeesmeet .	
12841	CGTCCGCGTT	TGTTTGTTTT	GGTTATTCTA	ATTCCACTO	TIGCATGCT /	LAAAGTTTAT
12901	TGGCAGTAGT	AGAATTTGAA	TTCTGGTTTT	TIGGTON CAR (	CIGATIGGT	GCATATTGG
			01031111	CIGGICACAT (	ATTAAGTGA 1	TAGTCAGTG

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12961	GAGAGGAC	AG GAAATCTGG	T TTATTTATT	ייידיתייינייינייינייי	T CCCCTCTCT	TGTTTGAAGA
13021	TGTTGATA	T CTCTGTGAG	G ACACAGGGT	T AGAGTTGGT	T GGGGTGTTT	TGTTTGAAGA TGACTTTACA
13081		W IGITITE	C TIGIATEC	ፐ ሮሞሞሞሮሮኒር	T TOO A A A A A A A A A A A A A A A A A	
13141	MOTOCHWW.	A GIIGICGAI	A TCTGCAAAA	ር ርክርጥአምኮርር	T CTCMMANA.	
13201		T GCCCIGIIA	I AACTTTTGA	מממממריידים כ	C TOTAL AND	
13261		n icurogual	CAAATGTCT	G GTCTC۵۵T8	A CTCCTNMCCC	
13321		W TIWWITTIN	G TAATTTCAC	A TITATITCOCO	C THE CAME A	
13381		w gwwgwwych	I AATGITGTT	A CAAATTGGA	ር ጥለጥጥርአርመርክ	
13441	AGAGTGCTT	T CAATATCTG	A ATAAAACAA	A GATTTAGA	T TTTCTAAACC	GGAAAAAAA
13501		G CHIGIGHIG	C TGGAAACTA	3 ርኔኔአሶጥአራአ	A THEORY COMMON	
13561	CAGAATTAT	T CATATTCTC	A GCAGTGGTG	C CACCTGACG	G ACTICITATE	AACTGAGAAT
13621		- CITIMACIG	A ICAACATEC	אידאכומידעעע יו	7 CCMXMCcccc	
13681	CACTTTAAA	T TCTGTTCTA	T TAGCACGGT	r Accorracer	A CCTATGGCTC A ATTGGCAATA	TGTTTTTACC
13741		* ********	A GACAGAATT	יי אינו הייירי אינו אינו	~ ~~~~~~~~	
13801	CACAATCTC	G GCTCACTGC	A ACCTCTGCC	r concerno	F AGCAATTTTC	GGTGCAGTGG
13861	CTCCCCAGT	A GCTGGGATT	A CAGGTGCAC	ACCACGCCT	GCTAATTTGT	CTGCCTCAGC
13921	TAGAGATGG	G GTTTCGCCA	T GTTGGCCAA	A CTGGTCTCG	A ACTCAGGTGA	GCATTTTAG
13981	CCTCCCAAA	G TGATGAGAT	I ACAGGCGTG	GCCACCGTC	CCAGAAAAGA	TCCACCTCGG
14041	TTATGAATT	T AAATAATTG	F GAAATTATCO	COCCACCOIG	TAAATAATA	CTATCTTATT
14101	TCTTAAATT	T TAGTTGGCT	F ACATAAAGAC	TOTANAGGG	ATTAATAAAT A TCAATTTAAA	TATAATGTAA
14161	TTTGTCTAA	A AAAAAATCA	ביים ביים ביים ביים ביים ביים ביים ביים	. TINNMATHCE	A TCAATTTAAA A ATGTGCTACC	TAAAAACTCA
14221	CTAATTAAG	A GAAAAAAAG	TTAACTGTG	GTTTCNTTN	TGGTCTTAGT	TCTTTAAGTT
14281	AAGTATTTT	TAAAAAAA	ACTTCACAAT	TTTTTATIAC	CTTAAAAATA	TAACAGCTTA
14341	TTTTATTAG	TITTTTTAAT	CAAGGAAAATA	יאראלייים מידער איייייייייייייייייייייייייייייייייייי	CTAATCAAGA	TTAATACCTC
14401	GACAAATTG	CTTAATAATT	TCATTTTAAA	AATGGCTTCT	TTATTCTTAT	TTATTTTTTG
14461	TAATATTAG	AGAATATTAT	AGTATACACA	DCLLLEDCCIICI	TCATATTCTAT	ACTGTAAAAA
14521	ACAAAAGCT	ATTTAACTTC	CATTTACTAA	LODDATITUM.	CTAGTTGTAC	AAAAACAAAA
14581	AGTTAACAT	ACTTTATTTA	TTATTCTAAA	אינים ברובר היים אינים אינ היים או אונים אינים	' ATTCATTGAA	TGGTTACATG
14641	TGATAATAGA	TAATGTCATT	TTTAAAAATG	CALIGIAAAII	TTATGTTACT	CCAAATTAAA
14701	ATTCAATGT	TGAGCTTAAG	TACTGAGTTC	ACAGTGTATG	ATAACTTTAA	AATTATAAGG
14761	O' O' CLEAN THE TANK TO	WWWIIGHGIN	AATTAATTCT	רא אירידידירים מי	A TO COMPANY A	3 3 TOTAL CO
14821	TIGONGGIA	CAAAATACAA	ATCACAAGAA	ACAGTGTAGT	א א מייים שייים מייים יי	T3 2 C3 TTTTTT
14881		GAATAACCAT	TGATAAACAG	ATAAGAGAAC	<b>ልጥልጥርልጥጥር</b> ር	COMPA CARAMAC
14941		TITCGCCACT	TTAGATTTGT	AAATCATGTA	CTCTATACCT	CECCCCCCC C
15001	HOUNCERIGE	AGGITTIGGA	TGACTGCCTC	TCTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	N TO COM N TO CO	
15061		TOTITAMGGGG	CTATGGTTAA	TCCDDDCDCC	ででできる へでへつかる	MG3 3 6m3 6m3
15121		CHALACACA	TAAGCATTCC	ひにりかりりがべって	MA COMMON OF	<del></del>
15181		GITACIGITI	GTTAATGTGG	TACATTCAAT	サイカ ヘヤス かへへる	MM 0 M 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0
15241			LICITATTT	TATATACCCA	יוו עידי עידידיים עיבורים	MAIMI MARK
15301	O TIMITANC	CWAICITCIC	TGACAACATT	<b>ልጥል አ</b> ሮአ አጥርር	TOOREGOOM	
15361	~ + * C-12-24 (************************************	CWWWINCIPC	ידידיי) מיזי ביו דו דו	CACACCACAM	773 M3 MARGO	
15421		GOWAICICAC	IGAGAAATAC	מידים מידים מידים	A A A A CO A CO .	mamaa
15481		CICCAGAAII	C. I.G. AAGT AG		7MM7111	<b></b>
15541		OWNERSHOW	GUTTUTTUTT	مادي لاملململيانان	CTCCTTTTTTT	
15601			LITCCAGITA	יייייייייייייייייייייייייייייייייייייי	TOMOGOGOGO	
15661			AATCTCAGTT	مان لامارششش لامل ك	3303M3mmoo.	<b></b>
15721		TAGAMMCAMG	GACCTTTATA	א איזיין אבויוייניים א	3300000000 ·	
15781		WITTWITT TO I	GLLTGTGGCC	ר אובי דידידי אין אין	TORRESSE.	
15841		GITIMACINA	AGTTGGCCAA	די ד	CACACOMAMM (	
15901		WILLTIGGW	GACAATTTGT	אבויים א מידיידים א	A TOTAL PROPERTY OF A	
15961	THE COLUMN	WCITITIE.	TGCCCTACTT		CORREMANA COR	
16021		WAGCCWIIGI	CAAAAAAAACA	ת ת ת ת מ מ מ מ מ מ	AACTEANAN .	~~~~
16081		TOCTITION	GATATTTTTA	<b>こころろろろろかべつ</b>	RCCROMMON -	
16141	GCCAGAAATC	GTGAAGACAT	GGCCTACCTA	ACTTGGAAAT	GTTGGTTGTC A	GTGGDDAAT
				·		

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16201	ACTACACA	GA GATAGCCA1	A GTGCTGCAC	A GCCAATCTT	'A AGTGTTTCT	A GAGAATCACT
16261	WHIIGHII(	- I AGAGAATCA	C TAATIGITT	מיאברדידים אי	ጥ ጥርጥጥር/ርጥጥጥ:	TACARGAGA
16321	GAGTATCCA	IT ACTAAACTC	T TTTCTACTG	A AAATAATGT	א מידית המממה הא	7 7 MOOM 7 MOO
16381	IAGACAGT.	IT GIAGITITI	T TCTCCCATT	Т СТАТТТТАТ	יתיתיטית מיטית מ A'	
16441	TIGITGAG	IG AAATCAGTC	C ATTGCTTGA	T ATACCTTGA	G CACAACTAA	TRACER MOCOCA
16501	AAAATTAAA	IT GICTITCAG	T CACAGTTTG	A CAAACTCAA	C TACCCTCACC	
16561	GIAAIAAT	G CCCTACTCA	I AAAGATGGG	G TGAAGATTA	A ATCANATACO	***********
16621	CACIAGITIC	C AGACGIGGI	A TCATGCTAG	T AAAATGGCT	G CACAGCACTC	CECTARCAGO
16681	ACAAAAAGI	G AAGCTTCTG	G AGACAGACT	C CAAGTTTGA	C TCCCACAMOR	CCACATGATG
16741	GATGTGGG#	C TCTGAGGCA	G GTCATTTAA	Т СТСТСТСТСТС	C ATTAGTATCO	CCACATATAA
16801	CTTTACAGI	G ATGGTAATA	G CACCTACCT	T CTAGAAGTA	T GTGAAGATTA	TICICIATAC
16861	ATGCATATA	A ACCACTGTG	T TTACTGCTG	T TTGACAAAT	T TTATTTATAA	AAGATCCTTA
16921	GCTCCTAAA	A GGACTTGAA	G CAGCTTATG	A CTGAAGACT	T TGGTAGGAGT	CCATCTTTAC
16981	TAAATTATA	A GAATTTCAT.	A AATTATTTG	TATGAAAAT	G CCAGTTGATC	TGGCCTTCTA
17041	TACCGGGGT	C CAACAGGTT	G AGAAAAAT	4 CACMMMM	CCCTGAACAT	ATAGTATGTT
17101	CTCTCTAGG	C ATATTCCTA	A GGACTTAAA	AATGATAAC	F ATCATTTCTC	ATGAAATTAG
17161	CAGATTTGG	A AGGATATAT	A TATTCAGCAC	ATTGACACA	ATCATTTCTC AATCCCAGTA	TTAAATCTTC
17221	AAAAGACAT	T AAAAATTAG	I GAAACTTTT	רדברטאטאט רדבררייייידיאי	CCTGTGTAAT	GTCCTAAATT
17281	CAAGCATAA	A ATTAAATTG	A GTAGAGTATZ	CINCETTIAN	ATTTCCTGAA	CCTGGATGAC
17341	GGCTCTGAG	T AATTTCTTT	GGGTCTGAAC	T DTCDCTGIAA	ATTTCCTGAA A CATATCCTCA	AGGTATTCTA
17401	GTTCATTAT	A ATTAAGAAA	AGGGAGTAA	TCTCCACAA	CATATCCTCA GAGCCACTTT	AGTATCATGA
17461	CTTGACCTC	A GTTCTTTTT	TCAGAGACAC	COTOTONO	GAGCCACTTT TGTTGCCCAG	CTTACTACTC
17521	TGGAGTGTA	G TGGCGCAAT	GCATCTCATT	GTANCCTCC	CCTTCTGGGC	GCTGCCAGGC
17581	CTCCTGCCT	C AGCATCCTGA	GTATCTGGAA	CCACACCICCA	TGCACACCAC	TGAAGCCATC
17641	TAATTTTTT	A AAAAGTTTTT	. ТСТАСАСАТС	CCACAGCAGG	ATGTTGCCCA	CATGCCAAGC
17701	AAACTCCTG	GCTTAAGTG	TCCTCCTCCC	TCACCCTAC	ATGTTGCCCA AAATTGTTGG	GGCTGGTCTC
17761	GTGAGTCAC	C GTACCCCGCC	' CCACTTCACT	TCAGCCICCC	AAATTGTTGG AAAAAATATG	GATTACTAGT
17821	GGGACTTTG	TTTGCTGATT	TABACATTCA	TCTGAGGAGG	TCATCCAATG	TAATAATAAT
17881	AGAATAATT	ATAGAGACAT	. CACCACACYA TYPYGYIICW	COMMONTO	TCATCCAATG	CGCAATTTGT
17941	GATCTCCTTC	CTGCTGGCTC	' ACAACCCTAA	GITTCTACAG	ATGATGGGGC	CTTGATAGTA
18001	CTATGAGGAZ	ATAGACCTAT	GTAGAGGGIAA	CTTOCHORGE	TAAAACCTTA	TTCTCTCATT
18061	TTAAAATTCT	AGGCTTATTC	TCTCACCATA	TCA A COMMO	TAAAACCTTA AAATGGTAAA	TCCTCATCAC
18121	TCAAGAGAAA	TATGAATAAA		CAGGITTIC	CCTCCTCTCC	AGAATTGGAT
18181	CCTTCCTTT	TTTTCTTCTC	CTTTTGTTT	TTTTCTC	TTTGTCTACT	CCCCATTCTC
18241	CAAACTCAAC	TGTAGGCTAG	AACAAAAAA	AATTICACTTT	TAAAATGTGC	ATTATTTGCC
18301	GTTAGACTTG	CTTABACAAT	ТССССТВ В ТС	AATTGAAAAT	TAAAATGTGC	CCCTTTTGTT
18361	CATTTGAGCT	TCAGTGCACT	CDADADADA	MACCTTGGAC	ACTAGATTTT AATTAAAAAA	AAAACACACA
18421	TGTTTAAAAA	ATCTGCAGAG	DARAIMAMIA	CEMERATE	AATTAAAAAA	TAAAATTGCA
18481	CTAGCCTCAA	GAGTGGATCA	AACAATACAC	GTTGTGAGAT	CTTGAATGGA	AGGAAAACTG
18541	GTTTAGAGAG	TGTGCTCAGG	GTTCTACCCT	GCAGGCAACA	GAGTAAGAGC GACAGTCCCC	ATGTTGGAGG
18601	CTTCGTCGCT	GTATCTTCTT	TATCABARAC	ACTARARATCA	TTTCCTCACT	ACGGCCTGGC
18661	TTATCCTTCA	AGTTTAGATC	AAATCCAAACT	ACIAAGICTT	GACTAGGTTA	GGATAAATTT
18721	TTAAGAGCGT	ACAGACATTC	PACCCCARCI	TAGGACACT	TTACTGCACA	CATTCATCTT
18781	CCAACAGCTG	TGCTACCTGG	CADACTTAAC	GGATGTGGGT	TTACTGCACA	GGCTCATTAT
18841	CGCAGGGAGA	ATGACAGTAG	GRANCI IAAC	CTCTCTGTGC	CTTAATTTCC	<b>CATCTATAA</b>
18901	TCTATTGTGT	AAAGTGCTTA	ANACACTCAIA	AGGTTGTTGG	AACAACTAAA	rgcattggta
18961	GCCATCATCA	TTATCATTCT	TOTORORO	TGGCACAGAG	CAAACATCCA (	STGAACTTTA
19021	AGAAGTGAAT	CAATCACTCT	CTCTCAGAGTC	AAATACAATA	TCTCATATCT (	SATAAATTAC
19081	TCTTTTCCAA	CAGTCGTCAC	TECTECTITTE	TCCAGGGGGA	GACAACAGCT 1	TTAGACATA
19141	TGAGTGATCC	TAGAAGAAGA	TAAATCCACC	TGTTTCATCT	TGCAAATAAA (	CAATGAAAA
19201	ACACCTGGCT	GAGAAAAATT	ACCTCTTTTTT	TATTTTGAAC	AATCAAAGAA (	GACAAATGA
19261	ATAGAAATTT	ATGACACAGG	AGCICITITIT	TCTATGCATA	AAACTATTAA A	LATATTCTTC
19321	ATTCTTTTTA	TATGTATATT	ATATATACAG	ACAAAATTAA	AAACTATTAA A	GTATCTCCT
19381	GTATCATATA	TAAAATAAAT	TTACCTORORS	MATTICATAT	AATAACTCCT F ATACATATAT C TTAGATAAAT A	TCACATCAT
			- INGGIGICA	IGATATATAT	TTAGATAAAT A	TACTTAGAA

19441	ACTTTTTT	ነጥ ርርስጥርጥአሞአ	3			
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19561	TACATAAA	C TICCCALLI	TATGCATTA	T ATTATAGAT	T ATATAGCTC	A CACATCTTTG
19621	AAATCAAA	C TITGIICAA	A TATTATTTC	C TAAGGATAG.	A CTTCATGAA	G TGGAAATACT
19681	CAGGATCA	TO TORMANACA	T TTTCTAAGG	T TCTTAACAT	A TACATTGCC	A AATTGCTATT
19741	TATTT ACA 2	T ARREST	A ATCCCAAAA	T AATATGAAA	A TTCCTGTTT:	T ATAGCACTCA
19801	*****	I WWWIIIIW	A AATCACTGT	T ልልሶሮሞልክሞክ/	_ TCCTTTC333;	
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19981	CACCTCCCT	T COCACTATA	T AAAGAATAA	ACCTTAGGT	r aaggccacat	TATCAATCAT TATTATATAAA
20041	0,10010001	T TICIGCGGA	G GACTCTGAAC	GGATACTAA1	א תיייניית ערובורים א	COCCIDEN
20101	CIGAAAIIA	C TITTACCTA	C ATTGTCTCTT	T ATAAACATTA	A TAACTACTCT	TTGAGAAAGT
20161	OTTINCIAL	G GACTGAATT	F TCTCCCCATC	CCCCCAAATI	איטרייית לות מידעם יו	700037733
20221	CCCMMINIG	A CICIATICE	L AGACAGGACT	TATAAGAGG	ושרים באת מיושר מעי	***************************************
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20641	CIMONCIN	G GAGAGAGA	AGTTAAACTT	'GTCCAAGGTC	<ul> <li>ACABABGCCX</li> </ul>	CARAGRAGE
20701	MADAGIGAM	G TIGACCTTGT	TCTCCTCAAT	CCAAGGCCAG	GACTCCTCCA	CTCCACATION
20761	HOMINGCCA	C CICACAGICA	ACAGCCAAAT	' GTCCACACCC	CAGAGTCAGC	ATTACACCA A
20701	GAIGICITA	CAGGAGACA	ATGCCTCATC	TTGAATAAAT	ATGTTCTAAC	A A CTTTA CCCA
20881	IGINAMACA	T TGAATCTCAT	: GAGAAACAAA	AATGCAAAGT	ATGTAGAAAA	CTATCTTTAL
20941	CUCTINACI	S ACAGTGATAA	AAAGCTTAAT	GATATCCTTA	TACTCTTCCA	CCCCmmmoms
21001	TAIGIGGIGA	A AACAGGTGCT	· CACGCACTGC	TGATAGACTG	TAAATTCCTC	CTDCDCDCD
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21121	ATACCI GGAL	ATTCGATTGG	CCATGCATCT	ATTTCTTCAA	TGGGTATGCA	CAGTTCACCT
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21301	TIMIGGAIAC	ATTGGGCCAC	ATTTACAGAA	ATTCACTTAC	AGTGGGTTAC	CACAACCCAT
21361	TATILITIES	GATTGGCAAG	AAGGCTAGGC	TGTTTTGTTG	GGGGCTGGCN	CCXCCTCTCT
21421	700C10CCC	AGTATGCAGG	TCTCTTCTAT	CATCCTGTGT	TAACCATCTT	CCATCTATION
21481	TICAMCCICA	TGGTCATCTG	CAGCATGTCT	AGGGGTCATA	TCTATCTTCC	ATCCACCA A
21541	AMAI DODAM	I GGGAAAGGGA	AGTAGGCATG	TACCATTTTA	ATCCACACCT	TOOMMONO
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21661	CONTRACTOR	CGICACCTTA	AATTCTAATG	TTTTTCTCTC	עטידיירטטידירט	A A A A CTC A CTC
21721	CNITAMOCTO	CACGTGGCTT	GGAAAAATTA	TTTCAGTCAT	CCAGTAATCA	CCTCTTCATA
21781	ABACAACATC	GGACATCAAG	TCTGTGTTGT	TAGCATTATA	CATGTTAAGC	ATTGAATAAA
21841	ASSECTATION OF THE PROPERTY OF	AIGIGGGTAC	ATTTCTTTAC	TTACATATAA	GTACTTATAT	A COMPANIA COM
21901	CHARDAGAG	GIIGAAAIGI	CAGGTGGAAC	ACABATARCA	TTROUTERAND	CEEE
21961	IGGGIGATII	TCAGCTATGC	TGATCTTTCT	TCTGGGTCAG	GTACTCCCAC	A A COMMOCOME A
22021		GCCCIGMICI	TAGTTCCTCT	CTCCTCTTAG	TC During DC N	COLOMBOSOS
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22441		CAGACAAATT	AAATACTCAG	TAGTTGAACC	**************************************	CONTRA S S M S S
22501	. OTT TOWN	MALGAAGAAA	TCATTTGAAC	י שמממדדשמאם י	י א איזייטיטיע א עדיין	**************************************
22561	OTHER PROPERTY.	TIGCCAATCA	AATATAAAGT	ፕሮኔኔኔኔኔፕነር י	TO COMPOSE A PAR .	
22621		GGACTACTCA	TTTTAAAAAAT	GTTAGATATC :	3663333666	2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2
	·	GTGCTGTCAA	GIGAAACCCT	GCTAATCTCA (	CTGAACATGT I	<b>AAAA</b> ATCTGT

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22681	AGATGCCT	TT ATTTTATT	ב כדראראכאכ	12		TA TGGTAAACAT
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22801	TATCACCG	GA GATAAGAA'	LL LYWWYITH	A TACTTTAA	A AATGGGCTG	T ATACTTTTCT
22861	CCATGACT	TT GCTACTTA	TALLALITY OF THE	T AAAATAAAG	T TATTTTCTC	T ATACTTTTCT T GTGACTGTTT
22921	AAATATTA'	TT TCAATAAT	ADADATION AS	T GCCAAAGTT	T ATCTAAGAA	T GTGACTGTTT A ATGTTTATGG
22981	CATGCCTG	TA ATCCCAGC	A RIGILIAGA	A GACTGAATT	T CCTGACTGG	A ATGTTTATGG G CACAGTGGCT
23041	TTCAAGAG	CA TOCTGGGC	CACAGAGA	C TGAAGAAGG	A GGATCGCTT	G CACAGTGGCT G AGTCCGGGAG
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24181			AGGCAAGGTC	רשידי עירו ביידייוייני		
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25201						
25261						
25321						
25381	GCCAATCTTA	CATAATGTGT	GAGATCTTGA	GACACCGGTC	ATGAGAGTGG	CCTGATTATG
25441	AGCTACAGGG	ACTTGGGAGC	ACCTOTANO	TATTACCCCA	TCCTTGAGAG	TCCTCTATAA
25501						
25561						
25621						
25681						
25741						
25801						
		TTTCCGCAGA	+ IAIICAGCA (	TAGACCCTG (	GAGATTCTG 1	TAAAGAGGGG

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25921	TTTTGTTAT	A CTCAACTTT	T CCGGGTAAA	A CAABCACAA	እ <b>ሞ</b> እርጥርርጥርር	CCAAGGGGCG
25981	GGGGCGGTG	C CTAGGTGAT	G CACCAATCA	AGCGCGCCC	T ACCCTATATE	A AGGCCCCGAG
26041	GCCGCCCGG	G TGTTTCATG	C TTTTCGCTG	G TTATTACAT	ר זיינכניתואן) ר זיינניניתיייתיי	CTGTTGTTAT
26101	GTCTGAAAC	C GTGCCTGCA	G CTTCTGCCA	G TGCTGGTCT	CCCCCTATC	AGAAACTTCC
26161	AACCAAGAA	G CGAGGGAGG	A AGCCGGCTG	CTTGATAAG	T GCDACTAIG(	AAGTGCCGAA
26221	CCTCTCTGT	G TCCAAGTTG	A TCACCGAGGG	CCTTTCAGE	TCACACACAA	GAGTAGGTAT
26281	GTCTTTGGT'	T GCGCTCAAG	A AGGCATTGG	CGCTGCTGG	TOACAGGAA(	GAGTAGGTAT G AGAAGAATAA
26341	CAGCCGCAT	C AAACTGTCC	C TCAAGAGCT	P AGTGAAGAA	COLLEGIA	G AGAAGAATAA G TGCAAACCAG
26401	GGGTACTGG'	T GCTTCCGGT	T COTTTAAGO	TAGIGAACAA	GGAATUCTGG	AATCTACCAG
26461	AAGCAAGGC'	T AAAAAGTCAG	TTTCTGCCA	GACCAAGAA	GIGATICCIA	CCAGGGACTC
26521	CAAGTCACC	A AAGACTGCT	A AAACCAATA	GACACCCAAGAA	2 DACCCOACAC	CCAGGGACTC CGACAACTCC
26581	TAAAACTGT	I AGGAGCGGG	A GAAAGGCTA	AGGAGCCAA	AAGCCGAGAG	AGCAGAACTCC AGCAGAAGAG
26641	CCCAGTGAA	GCAAGGGCT	CGAAGTCAAA	ATTENCECAN	CATCATCAL	TTAATGTTAG
26701	AAAGGCCACA	TCTAAGAAG	T AAAGAGCTTT	CCGGGAGGGG	A CATCATGAAG	GAACCCAAAG
26761	GCTCTTTTA	A GAGCCACCC	CATTATTT	ACATCCCCT	AAIIIGGAAA	GAACCCAAAG CAAGTTTCTG
26821	TGACAGTTAT	CTATAGGTTT	T ABGTTGTGAT	CCACCECACI	ACACTGGAAA	CAAGTTTCTG TGAGATTGGA
26881	GAATTAATTO	AGGCCAGGC	TCDDCDCCAT	COTOCOGA	TGAAAAGGCT	TGAGATTGGA TACCATCTAT
26941	ACCAGGGGT	CTCATTCCCC	CCCCCACCAL	CCCCTARAC	ATAGCCAGAC	TACCATCTAT ATGGCACGTT
27001	ATGAATTGAG	CCGCACAGCT	CAGGGGGTCAG	CCARCAMERA	GTCCCTGTCC	ATGGCACGTT TCCACCGCCT
27061	GTCAGGTTAG	CTGCAGCATT	· DEDTREBTTC	CGAACATTAA	CCAACTGAGC	TCCACCGCCT GTGAATGGCA
27121	CATGCAAGGG	ATCTAGGTTT	CAGGCTCCTT	CTCNCAAGCIC	AAACTGTATT	GTGAATGGCA GATCTGAGGT
27181	TGGAGCAGTT	TTAGTCCGGA	AATCATTGCT	CCCACCCCC	AATGCCTGAT	GATCTGAGGT GTCCGTGGTA
27241	TAATTGTCTT	ACACAAAACG	GTCTCTTGTG	TCAAAAAACCT	GCACCCCTG	TGGTTTTACA
27301	AAAAAGTAAA	TTAGTCAAGC	ATGGTTGGCA	CCCTCCCTTT	CTCCCTCC	TGGTTTTACA
27361	AAGGATACAG	TGAGCTATGA	TGGTGCTACC	TCACTCCAAA	GICCCIGCAC	GCGAGTCAGA
27421	CGTTGTCTCA	AAACTTAAAA	ADDIDOTACO	TTANARCACAGO	CIGGGTGACA	GCGAGTCAGA
27481	ACTGCCGTAT	ATCTAGAGGT	CCDGGDDCTD	1 1 MAMACAGA	MAAAGGGCTT	CTTGTCAGAG
27541	GATGGTGCAC	TAGAGGAGGC	TTTTACATETT	AAAAGICIGA	TGTCCAATCC	TGAAAAGCTC
27601	TCAGGGAAGG	GAAGTGGAGA	GCAATTTGGC	AMGAGCAICI	AAGTTCTGGA	AATGCCAGTG
27661	TTTTTTAACA	CAAGTACTAC	ATTOTAGTOT	TTCTGTGCTC	MACTIGUIGA MCAMMOMA A C	TACTTTTTT
27721	TAATATGCTA	TCCACTGACT	TCAAGGGATC	ATTANANTACC	AAMCAAC	TATTGTTTCT
27781	TGGATTAGGG	GAGTTTTTT	TTTGTTGTTG	THETHETHE	AATCAAGGIG	TCCCAGAATA
27841	GTAGCTGAAA	TTTAGAATTT	TCTTCCATTG	TGTGTGACTC	ATACARATA	TCATTATCCT
27901	GGTTATAGTT	GTTGCAAGAA	TOTGGAAATC	GTGCTTGCTT	ATAGAAATAA	CAAATTTGTA
27961	TATATCAACA	AAAACACACA	TATTACGGTC	ANGTOCTOCT	ATTICCGAAG	TACTATTAGG
28021	GGTCTAATAC	AATTGTAACC	CTATGAATTA	CTTTARCTRE	ATAATTATTT	TAATATTATT
28081	GTAAGTTTCA	TCAAACTACC	AGAGCATACC	CITTAMGIAI	CITATITATG	AAAAGAATCT
28141	TTAATGGAAA	TGTTGGAGGC	TGCCCAATTA	GGTTCTCDAR	MCCA COMMO	AATCCAAACC
28201	ACTTGTTTTA	ACTCTCAGTC	TGAGGTAAAC	TACGTTTCTC	TUCACUTICO	TGAATCACAA
28261	TTTTCCTTTG	ATTTTTGATT	TAGTATTCTT	ACTGATCATC	ATAAACAGA	LATAGTTTAA
28321	TTAGTCTACT	TTGGACCATG	GTATTTCGAG	AAACTTTGAA	CAAACTCCCC	MATGUTAATG
28381	TGCATTGCAT	TATTTCACAT	ACATTTATGT	TTTCCAGACG	CTTC3 ATS CT	1 GCAAAACTA
28441	TCTGAACTTA	TTTGTATAGT	TTGGCATCTT	TTTAAAAATT	GTGTCCTATAGI	ACCTUACTTT
28501	GTAAACATTA	TGTTTTAAAT	TTGTATAGAT	AAAATCAACC	ACAGACCTTTT	ATGAAAGGTT
28561	ATGTAATTGC	CATTGTTTCC	CAATGAGTTC	GGAATTACTA	CCATTCTCTC	CCTTGCTTGG
28621	TCACTTGCCT	GACATAGCAG	AGAGCCATTT	TGCCTAAATC	CTCTCCCCA	AAAATATGCC
28681	TCACCAGATT	CTCATCACAT	ACAGTGAGGA	TGAACAACTA	GCCTCTCCCA	CAATGGACTG
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28921	GONIGGAGCC	TUGUTUTGTC	ACCCAGGCTA	GAGTGCAGTG	GCACAATCTC (	CCTTC3 CTCC
28981	MCCICIGCC	TUCTGGGTTC	AAGCAGTTCT	CTGGCTCAGC	CTCCCACCTA (	COROCONOR
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29101	CTTGGCCAGG	CTGGTCTTGA	ACGCCAGACC	TCGTGATCCA	CCCACCTTCC (	CCLVCCVVVC STITCWCCWI
						CINCOMM

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29161	ACAGGGGTGA GCCACCGCGC	CGGACTTAG ACCACTTTCT TTTTCT
29221	CCATAGAACC CTCCGCAAAT G	ACACCTTCT CCCTARRORS
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29881	TITIGANG TICTCAGGAG AN	CCCCTNTT CTTN CACCA
29941	TOCOCCCTG ACGILLICAGE RE	TATATACMA CAMOCALAGA
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30061	CICGAGICIC CTCATAGATA AG	ACCCCAAA TCCCCCCCCC
30121	THE COCCUMENTAGE COUNTY OF A	GCCCTCCA TCTTATCCCC CA
30181	TAGCACCACC CITCCCCAAG PA	
30241	CAGIGGAAGG TATGAACTCA AN	
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31021		
31081	TOTAL COLUMN COL	TATAACAT TATTCOMORD
31141	CANADATAAA TAGAATAGCA GAA	
31201	- TATELY CHANGE CHA	CCTGTGG CACCTTTTCCA
31261	TIGITIANA AAACAAAAC AAC	CCCXCCC XCCXCmcm
31321	TINGGAMIAI TIAGAATEEE EM	CTCCCCT CTCACACACA
31381	TOCICACAIG AAGAGAAAAA ATA	CAGGAAT CCACAAAAA
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26347	GGCAACATAG CAAGTCTTCA TCTCTACTTA AAAA	AAAATA ACCAGAGGTG TTATGAAAT

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32401	ATAAATTG	C CAGAACTAC	C CTCCACAAA	C TAACTCTCT	C	G ATATGAGGAA
32461	TGAAATAT	G TGTGTGTGT	G TGTGTGTGT	G TATGTGTGT	C AGAATATIC	G TGTATGCACC
32521	TATATATG	C ACCTATATA	T TCAACAAAC	A ATTOTOATA	A TTCCCCARCO	TTGAGAATGA
32581	CTAGCAGC	C AGCATACAC	T ATCAGTTTT	A AGTATATA	T TOCCOMMON	TIGAGAATGA TAAAATGTAA
32641	AGAAATCC	A GAGTAGAAA	T ACTTTTAAG	C TATATTACA	G CTCACAAAA	GCATAAGTAT
32701	AGTCTCACC	C AACTTAGAC	T ATGGGGGCT	татаатаст. Т	A CARCAGRARY	GCATAAGTAT TTCCAGGCAT
32761	TTGGGGACA	T CACCACTGG	T CTTGGGCAA	G AAACTCCTC	T ACCORDAGE	TTCCAGGCAT TGATTTATCT
32821	CACTCCCAT	C TAAGGCTTC	A CTGCATTTC	T CTTTTTTCAC	AGCCAATGG(	TGATTTATCT ATTTAAAAAT
32881	ATCCATTTT	C TGATTCATT	T TTTTCTCAA'	T TANACTOTO	AACCTAACTT	CACACCTTTG
32941	GTTCCGTAG	C ATACCTGTG	T CTCTGCTGT	- 1999-01910)	A GIACCATIGO	CACACCTTTG TTACTTTTCT
33001	AGAAAAAA	T CTCTGCTTT	T TCTTTTCAG	CTITITIES	R CCTCCACTCC	TTACTTTTCT
33061	TTGCACTTC	C TAGGCTTGC	T GTCCTTGTG	F GGGCACGCTC	COMMANA	TATTAATACA
33121	CTTCGATTT	G TTAAAAATA	A AGATATOTO	COOCACGCIC	CCATAAACAC	TATTAATACA TTTTAAGATT
33181	TTAAAATTT	T TAATGTTTA	T TTTTTTCT	ACAGAAAAII	CAGTGGCACC	TTTTAAGATT
33241	ATGGTAGCC	T ACACTTCCC	C GGGCTCAAG	CATCCTCCC	CAGIGGCACC CCTCAGCCTC	ATGATGGCTC
33301	GGGACTACA	G GTGTGCACA	A CCACACCTG	. CTD ATTECTCC	TTATTTGTTT	CCAAGTAGCT
33361	TTGAGATGG	A GTTTCGCTC	TGTTGCCCAC	CIMAIIIIGI	AATGGCGGGA	GITTTGTTT
33421	CCGCAACCT	C TACCTCCCAC	GTTCDAGCAG	TTCTCCTCC	: AATGGCGGGA : TCAGCCTCCC	TCTCGGCTCA
33481	GATTACAGG	C ATGCATCAC	ACECCCACCA	A DETECTION	TCAGCCTCCC TTTTAGTAGA	GAGTAGCTGG
33541	CTCCATGTT	G AGGCTGGTC	LODRODOOR C	AAIIIIGIAI	ATCTGCCCGC	GACGGGGTTT
33601	CAAAGTGCT	G GGATTACAGO	CGTGAGCCAC	CACCICAGGIG	CACTAATTTT	CTCGGCCTCC
33661	TAGAGATGG	G CTTTCCCTG	GTTGTCCAGG	CACGCICGGC	ATTCCTGGGC	GTATATTTTG
33721	TGCCCACCT	r GTCCTCCCA	AATGCTAGG	CIGGICITGA	GAGCCACCAG	TTAAGTGATC
33781	AAAGATAAT	TCTAACATT2	מייייטיניטיניטיניטיניטיניטיניטיניטיניטינ	TIMULUGUGI MUUTUUTUU	GAGCCACCAG TCAAAAATTT	GTCTGGCTGG
33841	AGAGTAATTA	AATTTGATTT	י דים בסובונון	CTTCNATAC	TTCTTAATAG	TACAAACATG
33901	GCACAAAGTA	TTTTACATTT	י השתעתיים ארכים י השתעתיים או	TCANAMACT	AACCCAAACT	CACACAGAAA
33961	AAAACCGTAA	CATTATACCC	מממדדרם א	CACATCCCCT	CATATACATA	TACACAAAGA
34021	TGGGGGCAGT	AGTGAAGTTG	GTTATTTACT	GTTTTATCAA	AGTGCCATTC	GTAAAACTCT
34081	AGTGGCTCAT	GACTGTAATC	CCAGCACTTT	GCCACCTCCA	GGCAGGCTGA	AGCCGGGTGC
34141	AGGAGTTCA	GACCAGCCTG	ACCABABATGA	TGAAACCCTC	TCTCTACTAA	TCACGAGGTC
34201	ATTAGCTGGG	CGTGGTGGTG	TGTGCCTGTA	GTCCCACCTA	CTCAGGAGGC	AAATACAAAC
34261	GAATCGCTTG	AACCTGGGAG	GCGGAGATTG	CAGTGAGCCG	AGATCGCACC	TGGGGCAGGA
34321	AGCCTGGGAG	ACAGGGCGAG	CTCCGTCTCG	מממתממממ	ACAAAAAAGT	ACCGCACTCC
34381	TGACTTAGTT	TTAAGGAATA	AATCAAGGAT	ATTITA A CTICA	ATAGACTACA	GCCGTCATAG
34441	GTGACTTGCA	CTGAAAGTTA	TACGAATATT	CCTACTCA	CCCCTGCCCC	GTTAGCTAAC
34501	ATTAAAGACT	CCAAAATTCT	TTTTAGAATC	TTCACACTA	AAGCTAGAAT	TGAAGTATGA
34561	TAAATAATAA	AAAAATACTT	TGTATCTAAA	TOTCOTOTA	AAAATAACTT	TTGATTTTT
34621	GCTTCAAGGC	TATCCATCCC	CARATTTCTC	CCTGAATGAT	AAAGAGAATA	GGTGGATGAT
34681	TCAATTCAAA	AGTTAGAAAT	TTGGCCGGGC	ACGGTGGCTC	ACTCCTGATA	AATGAATATG
34741	ACGCTGAGGT	GGGTGGATCG	CATGAGCTCC	GGAGTTCAAG	ACCAACCTGG	ATCCTTTCGG
34801	CAGAMCCCCCI	TICAATAAAT	AATAGAAAA	AATCACCCAC	CCCTCCTCCT	
34861	CMGIMGGCIG	AGGTGGGAGG	ATCACTTGAG	CTCAGGAGGT	CCACACTCCA A	CTC N CCCCTC
34921	ATCGCAGTAC	TGCACACCAG	CCTTGGTGTC	AGACTGAGAC	CCTGTCTCAA	GIGAGCCGTG
34981	ACAAGTTAGA	AATTTGGCTG	GGCGCGGTAG	CTCACGCCTG	TAATCCCAGC	CAACAACAAA
35041	GCCAAAAAGG	GCGGATCATT	TGAGGTCAGG	AGTTCGAGAC	CAGCCTGGCC	ACTITIGGGAG
35101	AACTCCATCT	CTACTAAAAA	TACAAAAAA	CTTAGCCGTG	CAGCCTGGCC /	AACATGGTGA
35161	GTCTCAGCCA	CTTGGGAGGC	TGAGGCAGGA	AAATTGCTTG	AACCCAGGAG	IGCGCCTGTA
35221	CAGIGAGCCG	AGATCATGCC	ACTGCATTCC	AGCCTGGGTG	ATACACTORO I	CMCCTECEC
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35641	TOTAL GUARGARITA AATTAAGATT CAGAACACA COMARTON	
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37801		
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38341		
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38461		
38521		
38581		
38641		
38701		
38761		
38821	TCTGCCACCA CGCCCAGCTA ATTITTTGTA TTTTTATTAG AGATGGGGTT TCACCATGTT	
	NONTOGGIT TEACCATGIT	

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38881	NCCCN CCN	C cmc				
38941	TCCCATTO	G GTCTCGATC	T CCTGACCTC	G TGATCCACC	C GCTTTGGCC	T CCCAAAGTGC
39001	IGGGATIAC	A GGCGTGAGC	C ACCGTGCCC	G GCCTACTTC	A ביידייריריריריםי	מיאמממתים יו
39061	AATGGGGAT	A ATAGTACCT	A TCTCATAGA	A TTATTGTAA	G AAGTGCATG	7 3673376637
	GIAAGIAGG	T GCTCAGAAG	A GTCGGACAC	G AAGTAAGTG	ר איים איים איים איים איים איים איים איי	א מידיא מייני אייני אייני
39121	IIIICATTA	T CAGAACAAG	G AGAGACCAG	G TAGAAAATT	ייייייי מבוייבוריי מ	r TCACCTOMOO
39181	MAINCINGA	G TAGCATCCC	A AATGAAGGC	A CCATTAAAC'	די דייוכר א א דיריי	CTATCACAC
39241	TICATGCCA	A TTAGAAAAA	A CACCTCTTC	A CAACCCCTT	ד כממכמדמייייי	CCCTCCTT
39301	IGCIMMAM	C ACCCATCAT	A CTACCCACA	G ATAGCCATG	ր փնիրարդարարին	CCCACACAC
39361	CCICITCCA	T TCGTGCAGT	F TACAGCCTT	C ATAGCTGTGG	<sup>7</sup> አልሮሞሮክሮክሞሪ	ACREMONA COM
39421	GGAAGAATC	C CCAAGGCTT	G GTGACAGAT	G AGTTACTGGG	TABCACACAC	TONCON TONCO
39481	MAGGMAMG	I IGAAUGGGT	CAGAAAATG	C ATAGATACA1	יה ממממדברם יו	CTCCTTTTCCTT
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39601	GICCCCAA	A TITAAGGAG	CCTCTTCCA	A AAATAGGAAA	TGAAATGACA	TR COMORA DO
39661	TCTCTGAGG:	T GACGGAGGAI	ATGAAGGAAG	CCTCTAGATO	CAGCTTGAGA	TTCATGAGAG
39721	ACAGIICCA	<i>•</i> GGGAGAGGT(	ACAGCTAGGC	ATCACCGCC	TGCAGGAACT	CACAAAGG
39781	AATGGGGAAI	A TCTTTTTGAC	GAAATGAAC	GAGAAGGCTA	ADDTCDACCA	GTTCGTCAGG
39841	CAATTTCTAT	GTTTAGGTTC	AACTCTCTCC	TGAAACATGA	AGAGCTCATA	AATGCACTCC
39901	CTCTTTGAGT	CTCTAGTTTT	GTCTCCTTCC	CACAGTGAGT	CTGCAGGCTCAIA	CGTGTCACTC
39961	ACGTTCAGCT	AAGACGTAGT	GCCCCATGGC	TOTOTOTOTO	CIGCAGGCIG	ACCCAGGAAA
40021	GAGGCATCA	AAACCTAGGO	ACCATCTTGC	CTCTTCTCTC	TTCCTTATION	TCCTCATTCA
40081	CCCATCTCA	TTTAGACCTG	GGCACTATTO	GATTTCAACA	TICTIMITI	CTCATCTGGA
40141	AATGCTTATT	GGCTTTCTAA	CTGGTCTCCT	CVCCACAW	ACCATTATCT	CTCATCTGGA
40201	TCACCATATA	AGGGAGATCG	TGGTCCTCCT	TACETETEAL '	CIAACTTCTT	AACAACACAT
40261	ATCATAACCC	AATATCCCAA	AAGACCCCTTC	TICITAGGAT	CCTTCAATGA	CACCCCAGTG
40321	CTCTTTTCCC	TACACCACAG	ARGACCCIIG	GACICIGIAT	GAGCTGGCTT	CTTTCTGATT
40381	TAAGCAAACT	CAGGGTTAAG	GTACACTAAT	GGTAGAAATG	CATAATTGGT	GAGTGATAGC
40441	CTCCTACTTC	GCATGGTTGC	TCCCTCTCTC	TATITUTAAT	CTCCCAGTAT	GCCTTATACT
40501	ATGGAATCCA	GCTTCTCCTT	CARCATCICIO	AACCCTCCC	ATCATCTTCA	ACCTCACCTA
40561	TCATTCTTTC	GCTTCTCCTT	CANGAICCAG	AAGGCTATCT	TGATCCCCAG	CTGAATGTGA
40621	TCTTCTTTAA	CTACATTTAC	TTCCTATCAA	GCTTCCTGCC	TGCTTTAGGA	CCTCATGGGG
40681	AATAGCCACA	CTACATTTAC	ARCOMONNA	TTTCATTCCC	TACCAGATTT	GGGTTCTGAG
40741	AATAGTAGGT	GTGACTTCTC	TOTTO COMPANY	CCCCTGTACT	ACCTTAAACA	GCTCTTGCAA
40801	TATTTTCTCT	GCTCTGAAGA	TGTTTGTTGA	ATTAGAGACT	TTCATTCTGG	GGAGAACCAT
40861	TCCCATGGAT	CTCCCAGGGA	GCTGCTGGTG	TCCCCAAAGA	ATATAAATGA	GAAAAATGCT
40921	AAGAGACTTC	GCCAGATCCC	CTCTGCCCCT	CTTCCCACTG	TGCCCTGGGG	CAGAGGTACT
40981	AAAATTCCAA	CCCCTTGTTC	CTACTCACTT	GAACCCTGCC	TCTTCCTTAA	TATTATGAAC
41041	AGGGTGCCAC	TGAACAAGAT	GACGACAAAA	ACAGCAATTC	CACTGATGAC	TCCAATGACT
41101	CCCTCACCCAG	ACGGTGAGGG	CTCTAAAACA	GAAAAAGCAA	GTTAAAGCCT	TTGATTGCCA
41161	AGGCACTCCT	ACCCCCTAAC	AAAGAGCAGA	TCCTCATCTC	ACTGCCATAA	TTACCTCCTC
41221	CCCAGATCAC	CTCAACCCCC	AATAGATTTT	CTCAGCTCCT	GGCTCTCATC	AGTCACATAC
41281	CCTCTTTCCC	AATGAGGGC	TGATCCAGGC	CTGGGTGCTC	CACCTGGTAC	GTATATCTCT
41341	CCLATACCE	AGGGGGTACA	GCCAAGGTTA	TCCAGCCCTG	GTAGGTCCCA	TCCCCATTGG
41401	GCAMIACGIC	TTTAGGTTCG	AACTCCTTGG	CATCCATTGG	CTGCTTATCC	TTCACCCACT
41461		011616666	IAGTAGTTCA	AGGCCCGACA	CCGTAGAGTG	<b>CTC3 CTC3 3 C</b>
41521	MOGICACAIG	ATGTGTCACC	TTCACCAAAG	GAGGCACTTG	ACAGGAAAGA	GGBBGCBTCB
41581	GGAGAGGGGA	TCTGTTTACC	CTTGCCAGGA	AGACTGGAAC	TTTCECTTCC	がかいかみかれるこの
	TOGAGGAAGG	AAATACCCTT	TTCAGAAAAA	AACAAGCTAC	AGGAGAGACA	CCATTTTCTC
41641	ICCIMAGAIT.	GGACTCTAAC	ACAGTGTCAC	TTGGAGAGCA	GTCAGATCAG	
41701	ICACAIGIAA	ATATACATAT	CTGTTACCCA	TGTTCTTTGT	TCTGATAGAT	A A A TTCCCC
41761	TITALGIGCA	TIGAAAATGA	TTGAATACAG	ATGGTCAGTT	TCACCTGGGT	CDACCTAGGA
41821	GGCATTGTTA	TAAGAAGCGG	ACTTGTAAGA	TAGGTAGCTT	CAGTGATTAT	TGCTATGTTC
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41941	TITIGICCIG	GTATTCATAT	GATAGCTGAG	ATCTCTGAAT	արդարարարարարարարարարարարարարարարարարար	the bridge of the bride of the bridge of the bridge of the bridge of the bridge of the
42001	TITITAAGAT	GGAGTCTCAC	TCTGCTGCCT	AGGCTGGAGT	GCAGTGGCGC (	こみででででにここで
42061	CAGTGCAACT	TCCGCTTCCC	AGGTTCAAGC	GATGCTCCTG	CCTCAGCCTT	CCAATTAGCT

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42121	GGGACTACA	C CTCCCCA ma	3 omen			
42181	TCACCATCT	COMORCACO	A CTGTGACCA	G CTAATTTTT(	TATTTTTT	A GAGACGGGTT
42241	TCMCCWIGI	T GGICAGGCI	G GTCTCAAAC	I CCTGACCTT	* TCACCACCC	COMOCOCOMO
42301	TARCACCEA	.I GGGATTACA	G GGGTGAGCC	A CCGTGCCCG	CCTTGACATT	TCTGAATTTT
42361	TCCCTTTCA	G CCATATCA	A AAGATTATT	GTTAAATAA	AAGCAAGGG	CATAGACACT
42421	ACATOTON	T TATARCA	T GGAGAAAAG	AATTAAACC	ATGACTTGTO	GCTGTCTCAT
42481	AGGCACACA	C CCTCACCCA	G AGACTCTAGO	ATTGAGAAAC	TCCCTTCCC	GAATTTGGAG
42541	CTCCACACA	C CCACRACKA	C CTCTGAAAC1	CCAACCAGGG	ATTCCGTGCC	CTGCAACCTC
42601	DISTANCE	G CCACTAGAG	T ATAGGGGCAC	AAGTGTGTTT	CCACCATACO	TTGTTGGTCC
42661	TAGGCCCCTC	T TOTOCHAGOT	C CAGCAACTGO	TGCAGCTGTG	CAGGGCAGTC	CCTCTCCAGG
42721	GCCCTGGCT	T CTCCCTGG	C CCGAATCTTC	TGCCTTTCCC	ACTCCAGCTI	GGTGGGCCAG
42781	CCATCATAC	C CCTACTTC	r ccaatccag	GTGTCAGGGC	AGAATTCAAG	GTGGTCCTGC
42841	AGGATGACC	T GCACCCTCT	A GTAGCCCTCG	GTACTGTTGT	CTTCTTGCAT	TTCACAGCCC
42901	AGGAATAGG	T CCCTATTO	GGACTCTGGA	AAAATCCCCA	GCCTTGTTAA	CTGCAACCAA
42961	TTCCCTTCTT	CCCCATTIC	ACCATCCCCA	AGGACCAAAT	GATCTCAGGA	AGCAAATTCC
43021	AAAACATCA	D AACCTCCCA	A CAAGACCTCA	GACTTCCAGC	TGTTTCCTTC	AGCAAATTCC
43081	TGTGGTTGT	AAGCICIGA	AACCTCAGGA	AGGTGAGGCC	CCCTCTCCAC	ATACCCTTGC
43141	TCAGACTCT	S ATTITICATA	ATAGTCCAGA	AGTCAACAGT	GAACATGTGA	TCCCACCCTT
43201	TTCGGGGGCT	S ACTUAGUTGO	AGCCACATCT	GGCTTGAAAT	TCTACTGGAA	ACCCATGGAG
43261	-10000001	- CACACGGGG	L CICICATGAT	CATAGAACAC	CABCACCTCC	TORMOST SOF
43321	AGTGCAGAG	TTCAAACAAC	GAAAGACCAA	GGTCCTGCTC	TGAGGCACCC	ATGAAGAGGT
43381	GGAGCAGGA	A GTGTGAACCT	GGAGACAGAG	CAACAGGCCT	TAACCATGTG	TAGTAGGAGG
43441	CCTCATTTT	GTTGAGGCTC	CACACACCTG	CATCAACTCA	TACCATCAGC	TGTGTCTGGT
43501	CLCLLLLCCL	TGAAGGGTGA	GTTGCAGTCC	TGTCTTTCTT	CCATATGACA	GTCCTGGGTG
43561	TCCTATTCCT	GTGTGCTTTT	CTCTGCCACA	CGTGGCTGCC	ACCCCCTCAC	TGCCCCCAGA
43621	TTAAGGTGTG	ATACTCATGA	TTAGACAGAC	TCCACTAAAG	CTGGTGGATT	CTAGAAAATG
43681	CCAAATCCTC	TCTAGCCATG	GTAGTTGAAC	TCAGGAGTTG	GTGCTCAGGG	CAAATTAGAC
43741	GAGACAGAGT	AGGAATAATT	CUTTCAGTTT	TTTTTTTTT	TTTTTTTTT	TTTTTTTTT
43801	CAACCTGCAC	CTCACTCTAT	CACCCAGGCT	GGAGTGCAGT	GGCACAATCT	CAGCTCACTG
43861	TATAGGCGTG	CTCCTGGGTT	CAAGGGATTC	TCCTACCTAA	GCCTCCTGAA	AACCTGGGAC
43921	CCATGTTGGC	CGCCACCACA	TCARGCTAAT	TTTTGTATTT	TTAGTAGACA	TGGGGTTTCA
43981	CAAAGTGCTG	CAAGCTTGTC	ACTOROGOLO	GACCTCAAAT	GATCTACCTG	CCTCAGCCAC
44041	CTGAACTGCC	GGATTACAGA	CACCAC	CGTGCCCAGC	CTTGGTCCTG	AATTCTTACA
44101	TCCAAATGCA	TATGTGGCCT	TTTACCCCCTTGG	AAGCCTGACT	GGAATCTCAA	ACTTAACATG
44161	AAATGGCATT	GATCCTTGAT	CCACTCCCAAA	CIGCICTITC	CTCTGCCTTC	ACCATCTCAG
44221	CAACTTTAAC	GCCAATTACC	TTCACCCCCA	GGCCAATAAA	ATTAAAATAA	AGAACAAAGT
44281	AGGCTGAAGT	TCTTCTCTTT	TTCAGGGGGT	CAGGGGAGAC	AGGGTCTTGC	TCTGTCACCT
44341	AATACCCTCC	ACAGTGGCAC	CCCCACTAGG	CACTGCAGCC	TCAACTTCCT	GGGCTCAAGC
44401	GCTAATTTTT	ACCTCAGCCT	TAGAGAACC	IAGGATCACA	GGTGCATGCC	accacacca
44461	ACTCCTGAGC	GTATTTTTTG TCAGGAATCT	CCTCTCCTTC	CCCCCCCCCCC	GTTGCCCAGG	CTGGTCTTGA
44521	AGCCAATTCT	TCTCTTTCTC	TCDCDCDACA	TACARGOOM	TGGCATGAGC	TACTACACCC
44581	TATTCAGGAG	ACAATGGTTT	GTCACTCCCT	TAGAATCCTT	CAGCAACTTC	CTTCAGAATA
44641	CTCTTGCCTG	GACTGTGTAA	CAGCTTCCTG	CCTCCCCCCCC	CACCCAGCCC	ACTCCACTAC
44701	TTCATTCTGC	TTTCCACATA	GCAGCCAGAG	Cy y accommon	CIGCITITAC	TGTTGCTCCC
44761	GTTACTCCTT	GGCTAGAATT	CACACCACAG	COTTON COCC	AAAGCCTGTG	ACAGATCACT
44821	GCTCCTCTTC	TGAGCCCATT	ACCTACTTCT	TCCCCMCTT C	CCTGCACAAC (	CTTGTTTGTG
44881	TTTTTTTCAA	CCCGAGCTTC	TTAACCAGGA	CONTRACTOR	TROCTOR CAC	TACTTGTTTA
44941	TTAGAGACAT	TTTTGGTTGT	CAAGACTGGG	GENERACE :	INGGIGACAT (	GTGGCAAAGT
45001	THURST	CIGCINGACA	TCCTACATGC	AGATGGTAGT /	TOCCOTROCO .	
45061		CCCACACACA	CACACATGAG	<b>で及らすらですられる</b> :		
45121.		CCCMC TCMC T	TTGCCTGGGA	י הייהובייה מידממ	~~* *******	
45181	TTCCTTCATG	TCTCTGCTCA	AGTGTCAGCC	CLACACACACAC	CAGTCAATA 7	CATTCTTAT
45241		CONTRACTIC	CIGATGITGT	י ידירי דירידיידיירי דידי בעד ב	TOMORROSS -	
45301	CTCTCCCACT	AGAATGCAAA	ATATCANAGE	GTAAACRITCT (	TOTAL TERMINA	TATTGTCAT
				GINNIGACTT (	FITTUCCTGC 1	CTCTCCCTT

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45361	GGGGCTTC	3 C3CDCC333				
45421	TCTCCTCNN	C STORRE	A CATGGCTGG	G ACTCATTTA	ACTTGTAAA	C AATGAATATT
45481	AGCTATCAA	C TOGRAATTT	TATTATTCAA	C CTCTAATGC	A GTGTGATGT	T TAAGAATCAT
45541	CCCACCAA	G TGGAGACAT	G AGCTCTGCC	A CCAAAGCCC	GTGTACCAT	T GAATAAATTT
45601	accuracy will	C AGGCCGTGC	C ATGCCTCAT	I CTTGTCATG	Γ. GTAAAATCT/	CAMAGAGGMA
45661	GINCCHMM	C TCAAAGTGC	T GTGCTGAGG	C CGGCGTGTG2	CCCACACAA	T TOTOTOOMS O
45721	DUDAJAIJA	C AAAATCACT	G TCAACTAAG	A TTAGAAGCAC	CTGTAGTAC	T TC333T33C3
45781	TONGMANAC	C AGATTATT	A TGTTCTTTG	I AACCTGBBB	CACTTATATE	A TOTAL A TOTAL A
45841	CHGIINACI	I CIAGIAAAA	T AAACGTATT	<b>TTAGCTCCT</b>		7 0000 0000
45901	VICKWINK	G ATCAGATAT	G AATGTAACT	L AGAAGTGAGI	י כרש תתכניתייי	
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47461	GGATGGAGTT	TATGATAAAG	TCTGTGGCCA	AAATATCCTG	GAGADAGAGA	AACCACCCTAAG
47521	CAGGTGGAAA	TTCCCTAAGG	TGGCACATGC	CCAACAACAC	AAAAGCCTCT	AAGGAGGGTA
47581	ACCCCAAGTT	CATCATGCCA	TCATTATAAT	AGAATTTACA	TACACTOTO	COCCCCC
47641	CCTGGGAGGC	TTTTCTTAAC	AAATTATAGG	TAAGACCATG	CACACTUTAX	TTTTTACATE
47701	ININGCIAIA	AACTTCAATC	AAATAACATC	ATCCTGTCAC	TCAGATACAG	CCCABBCCMC
47761	AMCICCICCO	CACAAACCCC	ATAAAAGCAC	CTTGAGCTCT	GTABAGAACT	CCTCACTOCA
47821	CTTCGCAGAA	ATAAGCCCGC	TGTCCCTCAG	AGTGTATTAT	TGTGCTTC3 x	GCTGAGTTCA
47881	TTTAAGCTTG	CATTTTGGTG	TTAGTTTGTA	GTTCTTTCCT	CACTACACA	TAAACTTTGC
47941	TTGCTGCTTC	AGAGCTCCGG	CTATAATAAT	CTCCTCGCTT	DANGGATGGA	AGAACTGAGA
48001	TAATTCCCAG	TAACAGTATG	GGATGCCACC	TGGGCAATGG	CATHERINA NA N	TCCCAATGCA
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48481	CCAACATGGT	GAAACCCCGT	CTTTACTAAA	ZZTZCZZZZZZZ (	JGGTTCGAG	ACCAGCCTGG
48541	GCAACTGTAA	TCCCAGCTAC	ATTACACCOT	CDCCCDCCDC	TAGCCGGGC	GTGATAGCAG
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48601	CGGAGGTT	GC AGTGAGCTI	A CATCCTCC	1 magaz		
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48781	GAGATCATO	C CATTGCACT	C CACCEMOCA	T GAACCTGGG	A GGCGGAGGTT	GTGGTGAGCC
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49021	GACAATCT	TT AATCGCCIC	T CTTTCT	CATTCTCCT	CTCCTCCCCI	CCCAGCCCTA
49081	TATATTGCT	T TGCCGTGAC	T CCCTTCTATT	r GGAACATTT	A GTATACATAG	CCCAGCCCTA AGGCATATAA
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5196	AGGAGAATTG CTTGAACCCC COAGGAGGC 1G1AG1CCCA GCTACTCAGG AGGCTGAGGC
5202	CTCTAGCCTT GGTGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGA
5208	CTCTAGCCTT GGTGAGAGAG CAAGACTTGG TCTTAAAAAA GAGAAAAGAA AAATGAAATT  TCAGCATTAT AGAATAAAAA TGTTTCCCCT TCCCCCCAAA CTTTAAAAAA GCAGAAGTCT  GCATCATAAA ATGGTCTTTG CCAATGTTAT TTTTATTATA ACATAAAAA GCAGAAGTCT
5214	GCATCATAAA ATGGTCTTTC GGAATCTCTTC
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5226	TACCAGATCT CAGCAATTGT CACTATGTTC TGTAAAAATC ACTTCCTAAA ATGTCTGAAT TGACTGCTTG TCTCATTTAT TTGTTTCTCG TGTCATACTC CACTACCTAAA ATGTCTGAAT
5232	TGACTGCTTG TCTCATTTAT TTGTTTCTCG TGTCATACTG CAATGGATAT CTGTCTTGTT  AGTATAAATA TTTGTGCATT TTGTTGTTGT TAABACACCT TTTTTGTTGTTTTTTTTTTTTTTTTTTTTTTTT
5238:	AGTATAATA TTTGTGCATT TTGTTGTTGT TAAAACAGCT TTTTTTGGCCT GTCTTCTCC  ACCTATGAGG TAATATAAAA CTCATGTTTA ACACTTATTT TTTTTTGGCCT GTCTTCTCC
5244	GACAAAACCC CTCAGACACT GACTAGAGAGAGACTAGATTATTATTATTATTAGGAGG ACAAGCTAGA
52501	GACAAAACCC CTCAGACACT GAGTTAAAGA AGGAAGGGCT TTATTCAGCT GGGAGCTTTG GCAAGACTCA CATCTCCAAA AACCGAGCTC CCTGAGTGAG CAATTCCTGT CCCTTTTAAG GGCTTGCAAC TCTAAGGGGG TCTGTGTGAG AGGGTCATCA TCCATCTCTGT CCCTTTTAAG
52561	GGCTTGCAAC TCTAAGGGGG TCTGTGTGAG AGGGTCATGA TCGACTGAGC AAGTGGGGGT ATGTGACTGG CAGCTGCATG CACCAGTAAT CAGAACACAA CAGCAGGAGC AAGTGGGGGGT
52621	ATGTGACTGG CAGCTGCATC CACCTGTT CACCTGAGC AAGTGCGGCTT
52681	ATGTGACTGG CAGCTGCATG CACCAGTAAT CAGAACAGAA
52741	AACCAGACC AGGETGGAAC CTATAGATAA CATAACCGGT TAGGTCGGGG GTCAATCTTTT
52801	AACCAGACCC AGGGTGCAAC ACCAGGCTGT CTGCCTGTGG ATTTCATTTC
52861	CTTTTACTTT TTCTTTCTTT GGAGGCAAAA ATTGGGCATA AGACAATATG AGGGGTGGTC GCCTCACTTA TTCACCCCCT TTGAGAATCT CACTCATTAG
52921	GCCTCACTTA TTCACCCCCT TTGAGAATCT CACTCATTAG TGGGAGTTCT CACTTTTATT  CTCACTACCT ATGTCTTCTT GAAAGACAGA TTGATAATCA TGGGAGTTCT CACTTTTATT
52981	CTCACTACCT ATGTCTTCTT GAAAGACAGA TTGATAATGA TTCATATAGT ACACTTTTATT TGAAGCATTT TGGTGAGCTA AGGTAGTGAT GAAGCTTTTTT
53041	TGAAGCATTT TGGTGAGCTA AGGTAGTGAT GAAGCTTTTT ATCATTTGGA GAAGTACAGG TAGCAAACAA GGAAGCAGTA AGCAGGTTTC TATTAATATT ATCATTTGGA GAAGTACAGG
53101	TAGCAAACAA GGAAGCAGTA AGGAGGTTTC TATTAATATT ATAACTCCTA TTATAAGAGT TTTAAATCTT CTTAGCACTC GGAACCATTT TTCAAACATC
53161	TTTAAATCTT CTTAGCACTC GGAACCATTT TTCAAACATG GCCCCAGAAA CAAATCCATA
53221	CCACACCTAC ATGGGCACAT GTGCCACTTT TTCAAACATG GCCCCAGAAA CAAATCCATA TTGCCCTTAA TCATCTATGT GTAGACAGCA ATTACTAACTATGT CTTCAACTAC
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53401	CTGAGTTTCT TGCCAGGCCA CAGTAGTCAG GGCTCTGCTG GTCTTATTAG TAATTATTTC TAAGACAGCT TGTAACCGTA TGATTCAGTT GAGCATGTTA
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53641	TTGCGGGAAG CATATACAGG GAAGCCCAGG AGTTTGCCTG TCTTTATGGG CAGTAGGAAG AAAGATGGTT TAATAGTGTC AATAACACAA CTACCTGGGG TCTTTATGGG CAGTAGGAAG
53701	TAAGCTGTAT GCCCACATAT GGCCACATATT GGCA
53761	TAAGCTGTAT GCCCACATAT CCAGTATAAT CCAGTGGGGG CTGTCCAGTC CCGGTGGGAC TCTGGGTGGG TCCACACAGT TTGCAACTTT GCGAATTTTGGCA
53821	TGGTTTGAAC TCCACTAGGT CCCTCTTTTTTAGTG
53881	CAGCTGAGTC TTCCCACAGG AACGCTGAACATT TATACAGTTT TTGCCCAAGG
53941	CTAATGATTG AGGCTTTTAG CACCOAGAAG TCTTTCCCCA CTTTTGCTAT ACAGTATTGT
54001	TTATCAGGAA CTGGGTCTGT ACCTAGGAAT
54061	ATTACAGTTC CTCCACATAC ATACATACA
54121	CTCAGCTAAT TGCAAAAACA AAGTACA TGAAGAGACT GGGCTACATG
54181	CAGTTCATCA TAAGAAGGTT TOTAL
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54301	CCCCTTTTT CCAGTGAGAA MGAAGAGAAC TATTTCTAAG GTTACACGAT
54361	GACCACTEGT ACAGGAAGG GGGTTACACT
54421	TITTAACCA AGTTGCCTAA AMGAGA CAGA GIGGACAGGA TTCTTTTTAT
54481	TTAATTCATG ACAAGCGTAC TTATTCATTCATCATCATCATCATCATCATCATCATCATC
54541	CACATCCTAT TTCTAACTTA TTACTATTACTATTACCC TCTTTCCTAA TGAACAGAAC
54601	CTTGTTTGGG CATTCCTTTTT MCTTGTCAAGCTGA
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77021	ACTGGAGCAG GGCTTGTTGT CTTCTTCAGT CACTTTGCAG GCGTTGGCGA AGCTGCCACG
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55081	TACAGCTC	AC AGTCTACTO	A TGTTCAAGG	A TGGTCTTGG	A AGTTGGGCC	ACTAGAATTA
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55381	TAGCAGGG	T GTTGCTGCC	A AGGCTCTTA	A GCATGGAGG	ር ርልልርርርምሞአር	7770000000
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55501	CCATTTTT	C TCTTTCTGA	C ACATAGAGT	G TAAAGGGTT	т тетелестел	GGTAGCCCCA
55561	GGGCTGGGG	C CGACATGAG	T TTTTCTTTT	A ACTUATGAN	A AACTCATTGC	GGTAGCCCCA
55621	AATAGATGT	A GTTTATCCA	A TCTACATTT	T TATTAACTC	T CACCCACCAA	TGTTGGTTGT
55681	CHARTCCTG	C AGCTATTTG	A TTTTGGGAT	T TAAATTCAT	TO THE STREET	mamaaaa
55741	CAATTGCAT	C TAAATAGAT	G TGAGAGTTG	A AAGACACAT	A AGGGTCTTCT	TGTGGGACTC
55801	GATGTCTTA	T TTTTCCTCC	C TCTGGTTGA	T GAAATGCTA	G GGTGAAAGGG	CTTGCTTTAC
55861	GGACTAAAG	T ACAAGTGCC	G CTCCAGTTA	T TTGGCAGAG	golgaaaggg Goocagtaaa	ATAGCCAACT
55921	AATACCACC	A CACATCCGC	T TGGGGATGA	A CABAGGCTG	GCCCAGTAAA A CTGATTGAGA	GGTCCACCAC
55981	AATTCTTAA	G CTCACTGCA	CCCTTCAGG	P CTCCAAGGA	A TGCTAAGTTT	AGCTCCTGAA
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56161	CAGGCTGTA	G CATCCTGGA	A AACAGTTAC	T ATCCACACAC	TGCCTGGTCA	CTTATTACTC
56221	CACCTTAGT	G GAAAGGGGA	L AATCTGGCC	TOTOCOCOTO	CATGTGCACA	ACAGGAGGAC
56281	TTGGTTTTG	TTAATGTGTG	GACAGAATA	TOTOGCCIGC	CATGTGCACA CCAACTGGGC	AGCATAACAA
56341	TGGTATCCT	G CTTAATTATO	AAAGTTTGT	TIGATECAT	AACTTCTATG	ATTTGCATCT
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56461	TTTAGTGGT	CACACAACAT	TCGACCAACT	TATGGGTTANA	AGCTCTACAT	ATCCTCGCTC
56521	GACTCCTCG	TGACACTGG	GTCTTTATTG	AIGGCAIAAA	GGATTAAATG	CAGGGGGCAA
56581	ACTAAGGCT	AGTCTGAGGA	GAGTCAGGAG	CCACACACACCT	GGATTAAATG ACTTTTCTGA	GTCTCAGTTT
56641	TGTCTTCGA	TTGGCAAGTC	CCCACAGGGT	OGACAGAGGI	ACTITICIGA	AGTACAGAGA
56701	TGAGGCAAA	TTGACTTGGT	מידע מידים דבר '	ATAACAAGGC	CAGAAATAGA	TTCAATAGTT
56761	AAGAAAGAGT	AATAGAATAG	ATGARGACT	TARATEGI	TTAGCTTTAG	GTGAGGGAAG
56821	TTTTCCCCTG	GGACTATGGC	CCATGACTCT	GGACCCCCCCC	GCACTTTCTT	TTTGGTAGGG
56881	TGATGAGTCC	ATCCCTTTTT	Carronerer	ACAACACTC	CGGTGGTTAG	GACTCGGGTG
56941	TAGGGTCCTT	CCTAGGCTGG	CTCAACTTTT	CCTTCTTCT	ACCCTTTGAT	CAGCACAAGG
57001	TCTTCAGGCT	GGTGCTGGTT	TACAGAAAAT	TOTACCOCTC	GTACATGTGC	GAGAACATGA
57061	TTAGTTTTGA	GGGAAAGGAA	AGTGGAAGAT	D D D C C D D C C D	TATAACTTTT	TAAAAGACTT
57121	CCTTTTGTTT	TAAATGTGGG	GACATCAGCA	CTCCTCTTTT	TAGTCCTTGG	AAGAAGTTGA
57181	CTGAGAAATT	TCCTTTAGCA	CCTATTTTT	TTA CTTTTTA	GACCAAAGAA	TGCCTTCTTA
57241	CATTTTATAT	TTGACAACGC	TTCTTCTATC	TIMGITITIA	ATAAGCTAGA	AGTCAAATGC
57301	ATATTGGTGT	GTTATTAATG	TTABACTTAC	TITATACCAG	ATAAGCTAGA ACTCTGTAGA	TTTCACCTTT
57361	TGATTTTTAA	TGTCTGACCA	TAAGGTAAGA	TITIAMIAAA	CTTTTCTTTA	CATATTTATT
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57481	TTCAGTTCAC	AGAAAAACTG	TATGATACCC	CONTRACCO	GCCAATATGT	PATTITAATG
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6840	TTATGGGGTC TTGTTTGAGG TOTGTGTGT
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69301	AGGCGGCAG ATCACCTGAG GTCAGGAATT CGATACCAGG CTGGCTAACA TGGCGACCCC GTCTCTATCA AAAATGTAAA AGTTAGCCAG GTCTCTGCTGACA TGGCGACCCC
69361	GTCTCTATCA AAAATGTAAA AGTTAGCCAG GTGTGGTGGC TCGCACCTGT GGCCCCAGCT ACTCAGGAGG CTGAGGCAGG AGGATCGTTT GACCCGGGCT TCGCACCTGT GGCCCCAGCT
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74221	AAGGAGGAAG CAGAATCACA ATTAGGTCAA AGGAAGATAC GGGAGAATAA AATATGTATT IGGTCAGGGA AAGGATGTAT ACTGGAAGAG GAAGGGAAAA TAGAATATAA AATATGTATT
74281	TGGTCAGGGA AAGGATGTAT ACTGGAAGAG GAAGGGAAAA TCAGATATA AATATGTATT TGACTTATTA GGCAATACAA TAATAACTTT TAGGGTCATT TTTTTTTTAA AGTTGTTTAA
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74521						AAGATTTATC
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74641						TTGTTTCTTT
74701		AATGTGACCA				
74761		GTATTGCCTT				
74821						AGTTTTGCAT
74881						TTTCTCTGTC
74941						CTGGATGTCC
75001		AGGTCTGGAC				
75061						TCGATTTATA
75121						AGACGTTTCA
75181						GTTCATTGCA
75241		GATCTAAGTG				
75301		CCCAGGCTGG				
75361		TTCTGGAGTG				
75421		ATTCTCCTGC				
75481		TAATTTTTGT				
75541		AACTCCTGAC				
75601		TGAGCCACCA				
75661		TTCGACTGAG				
75721		GCAACCTCTG				
75781		CCCCAGCTAA				
75841		CTCAAACTCC				
75901	GGATTACAGG	CGTGGGCCAC	GGGGCCCAGC	CTTATATTAT	TTCTTTTACT	ACAATATATT
75961		AGGTGCTTCA				
76021		CTGAGGAATA				
76081		ATTAGACTGT				
76141		GACAAATGTT				
76201	TTTCACTTTT	CATTTGTCAT	TGAACTCTTA	TTTGTAGGTT	CCCTTTTGAC	TTTCCCACAA
76261	TCTAAGGCTG	TTCTCTTTAA	CACATATTTT	CATGAAAACA	TATATTTGAG	CAGAAATTGT
76321	TGGGGAGTTG	TAATATTACC	TTTGTCCCTA	AATATGAATC	TATAATTATA	TCAAATATAT
76381		TTTACTTTGC				
76441		AAGAAGTAGT				
76501		AAGAGGTGAT				
76561		TTTTCTTGTC				
76621		TCATCTTAAA				
76681		GATATATTTG				
76741		ACTGAAAGTA				
76801		AATAAATAA				
76861		GCCAAATCTA				
76921		ATCTTGAGGG				
76981		AAAGAGCCCA				
77041	CATCTATGAA	AAGAGGGATT	TGATAGTTTC	AATGTCTTCA	AATCAAAGAT	TTAAGTCTGT
77101		CACCCCGGAC				
77161		TGGCCGTGGA				
77221	GAATTCACAG	AGGACCTGTG	TTACTTCCCT	TGTGAAGAAA	CAGAATTATC	ATGAAAATTT
77281	AGGTGGAAAC	CATTTCGCTT	TTTTCTTCAA	AAATAAGGGA	AGCATGTGCC	CAACCACCCC
77341		AACCTTCAGG				
77401		ACTGCCCCAG				
77461		CGTTTGTGGA				
77521		CCAAACCCAA				
77581	TGAGGTGGAA	CAAACACAAC	TTGGGAGCAG	CGCAGCGGCT	CAGAGCCTGC	CAGCCAGGCG
77641	GGCGACCAGA	GCACCAATCA	GAGCGCGCCT	GCGCTCTATA	TATACAGCGG	CCCTGCCCAG
77701	ACGCTGCTTC	ATCGGCGCTT	TGCCACTTGT	ACCCGAGTTT	TIGATICICA	ACATGTCCGA

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77763	<b>~~</b>					
77761		CT GCCGCTCC	CG CTGCCGCGC	C TCCTGCGGA	G AAGGCCCCTC	TAAAGAAGAA
77821			JO GIACGCCT	G Transceceme	C COMPOSS	
77881		00101000	-G CCILIAAAI	A GEOGRAPHICA	A COMMONANT	
77941		TO OCTOCE	-G GLIATGATG	п ссастатата	~ ~~~~~~~	
78001		OC CIGGIGMG(	-A AGGGCACTC	ፐ ርርጥርሮአአአአሶ	~ ~~~~~~~~~~	
78061		un crevitati	M AGGCAGCCT	C CGGGGGN NCC	7 330000	
78121		CC VINTECTIMAL	A AGCCAGTTG	G GGCDGCCDA	~ ~~~~~~~~	
78181		or convolution	M GLGCIAAGA	<b>A AACACCCAA</b>	~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	
78241		er accusoum	LO TOCCIANCA	G CCCDABGBB	~ ~~~**	
78301	******	ar unantalet	G CTAAGGGTT	r <i>Candeeco</i> nn <i>i</i>		
78361		TO MAGGGGGG	IC CCHAGAAGA.	u ልጥአ <i>ርርርር</i> ር	~ ^^~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
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78541		** ************************************	O GGIII-L-L-AL-AL	. 1112222224	* ^**^*	
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78901	TATATTAAT	G TCTGGGATT	I CGGACGCTTT	CCATCIIANA	GGTAGTCAAG	CTTTTCTCTT
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79081	CCACAACAG	C GGCAATAGC	CTTCCTCCAC	CCDDCCCDAM	CGTGGACCTA	TAAAAAACAG
79141	TGTGCCACA	T AACATGTAG	CTTCCGCTAA	ACTCA CA CCT	TTGAGCGTAT	GGGAGTTTTT
79201	CGTATCGAA	A GCACAACTT	TAGCCAGCCA	TTTTCTCTC	GCATGACTAC	CGATTTTGAG
79261	CCTGTTTAG	A CAGACAGCAZ	אממממדדדאם ג	TCCAACTTCC	TTTAAACGTA	GGTTGCTTAT
79321	CAGTCCAAA'	T GTTTCTATGO	AGADACACT	ATTTCTA CON	TTAAACGTA	TTTTGTTTGG
79381	GATAAATGG	G AGACATTTCT	AATAAAGGCC	TTCCTTAATC	GTTCCCTCTG	AGAGTGTATG
79441	ATGGTGCTT	TGAATACAGA	AAGCCTAGCG	TOTTATATE	GCTTCTTTTA	TTTGACATCC
79501	GGCACATTT	r GGTGAGACCT	AAATTATGGG	CACTCCCCCC	TCTGGAGATA	AAATCTGGTG
79561	TTATTCTAC	ATCTCCACAA	ייאייה איייה א	ACTCACOTOCA	TTTGTTAGTG	AGCTGCTCAA
79621	CGGATTCAT	CCAAGAAAGA	GAAAGGGGAG	GCACCCAACC	AGAGAGACAG	ATAGTGACCA
79681	GCAGGGAAGA	AGGAGAAAAC	ATTOTOCOM	CCTTTTA	AGAGAGACAG ATTTTGTGTT	GAAGACAGAG
79741	CATTACAACA	CGGTTTAACA	TGGTGAACCC	TOTATEMEN	TGTAAGGTTT	GTTAATTTTA
79801	CATATTTTTC	CCAAGACCAT	TTATCAACTC	TCIATTTIGG	TGTAAGGTTT TTCCCCCCTTC	AACATATGGA
79861	GCCACCCTCC	ACGCTCCTAT	CAATTTTCCC	TCATTTCTGC	TAGGCTAATA	TTCCTCCCGT
79921	TCATGGACAG	TTGGACTGTC	TTAGGTTTCT	Caccommena	TAGGCTAATA	CGCTATAATT
79981	CCACAATTCT	TAAGGTAGAA	TTGTATTGTT	CAGGITTCTA	TGTTGTTCCT	<b>TTAGTCATTC</b>
80041	GCTGAGATGA	TTATGTGACA	AATGGCAAGT	CTTCAACATTG	TACCTAAATC	FATCCTCAAT
80101						
80161			LATCACARACT	7777777778*	~ ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	
80221	AGTCATATAA	TTATATTTAT	איייייים איייים אייים	TTATTTATA	GCTATTTTGA ( GACGGAGTTT (	BAGAGATCAC
80281	CCCAGGCTGG	AGTGCTGTGG	CACGTTCTCC	CCTCTCTCTCC	GACGGAGTTT ( ACCTCCGCCT (	CCTCTGTCA
80341	AGCGATTCTC	CTGCCTCCGC	CTCCCGACTA	GCTCACTGCA	ACCTCCGCCT ( CAGGGGCCTG (	CACGGGTTCA
80401	CGGCTAATTT	TTGTATTTTT	AGTAGAGACC	GCIGAGATTA	CAGGGGCCTG C AAGTTGGCCA G	CACCATGCC
80461	GAACTCCTGA	CCTCAGGTTA	TCCGCCCACC	GGGTTTCACT	AAGTTGGCCA G AAAGTGCTTA G	GCTGGTCTC
80521	GTGAACCACC	GTTCACAGAC	TCAAATCATT	TCATCCTGCC	AAAGTGCTTA G	ATTACAGGC
80581					AAAGTGCTTA G FATATTGTTA T FATAAATTAA A	
80641						
80701	CATCCACTGG	GGGTGCAGTT	TATTALATA	CCDUMUS CO	STACTATCTG T PAGTCTCCCC T	GGTTTCAGG
80761	TAATTAACTG	AGATGTTGTA	ACGTGACTO	GCATTTACAT	PAGTCTCCCC T AGAGCTAATT T	TTGGGAGAC
80821	ACTCTTCTTT	TTCAGAATTT	TCCTCCTTATE	MATAGCAGAT I	AGAGCTAATT T ATTTTTCCAT A	TCTCTCATT
80881	AGATCTCTTC	CACCTCCTCC	TGTTTCTCA	TCTCARTTTTT )	ATTTTTCCAT A AACAATTAA A	TGTATATTA
80941	AAAGGCTGGG	CGCGGTGGCT	CACGCCTATA	ATCCCARCATC	AACAATTAA A TTTGGGAGGC C	AAAAAAAA
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81001	GGATCACC	NC CECTOS				
81061	TAAAACTX	MG GICAGGAG	TT CAAGACCAG	C CTCGCCAAG	A TGGTGAAAT	C CCGTCTCTAC
81121						
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_		-CALLITAAG	ATTAGGTGTA 1	CTGCCTGGT 1	CTCAATTTG A	CACCCTTTC

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04343	MGMGma a a co	<b></b>				
84241	TCTCTAAAC	A TGAATGAGT	CCAATCATA1	TTATTCCTA	GCTATCACAC	TCAAATATAC
84301	TACAGATCT	<i>i</i> TGGAATATGO	: CAAAAGTTAJ	L GGTGAAAAA1	፣ ጥልልልጥተልጥተ፣	CCTATTTTCAT
84361	AGTTTTGCT	A GTTTTTGAT	TGTGAGTGA	TATAACTATO	CTCTATGTCC	TGGCACTGTT
84421	CCTCAGAAA	C ATAGGGTCC	CATATGTAA7	TTTAAATTT	די ד	א א א איזיייזיייזיייזיי א א
84481	AGTGGAAAA	A GAAATCTATI	TTAATGATT	GAATCCAGTO	ממממסטמאד:	א ערטיזייזייזייזייזייזייזי
84541	AAGGTATCT	ATATTAAAAT	ATTGAGTTT	TACTTTGTT	TTTTACTACC	ייי א א מיייייייייייייייייייייייייייייי
84601	CIGGIGIGIA	A TITTACACTI	. AAAGCACATO	: ACAGTTTGGA	GTAGCCACAT	TTCCNATCCT
84661	TAATACTCAC	C ATATGGTTAG	TGGCAACTAT	CTTGGACAGO	ACAGCTTTTA	TROTOTOCOR
84721	AGACACAAGC	CAAATACTTGC	: TCTGCAGCAG	AATCCAGATC	TTTTCCAACA	777C7C7
84781	TCTGACCTGT	TCGTGAAACC	CAGGTAGTGT	' CTCTAATACI	VILLAL TO WILL Y	THE CHATTER OF C
84841	CIATTGTAAC	CACCCAACGG	GCTCTCCTTG	TCCACTTCCT	' AGACAGAGCT	CDTTTTTTTT
84901	GACAGGGGA	I TTGCAATAAG	GAGCCAGCGC	TACAGGAGAC	TAGAGTTTTA	THENTHACTOR
84961	AATCAGTCTC	CTTGAGAATT	' TGGGGACCAA	AGTTTTTAAG	GATAATTTCA	TTCTACCCCA
85021	CCAGTGAGTC	GGGAGTGCTG	CTTGGTTGGG	TCAGAGATGA	AATTATAGGG	ACCCTAACCT
85081	GICCICTTGI	GCTAAATCAG	TTCCTGGGAG	TGGTGGGGTG	GGGGACTCAA	GACCACATAA
85141	TCCAGTTTAT	CTATATGGGT	' GGTGCCAGCT	AATCCATTGT	GTTCAGGGTC	TGCAAAATAG
85201	CTCAAGCATT	GATCTTAGGT	' TTTAAAATAG	TGATTTTATC	CCCAGGAGCA	ATTTCACCTT
85261	TAGAATCTTG	TAGCTTCCAG	CTGCATGACT	CCTAAACCAT	AATTTATAAT	CTTCTCCCTA
85321	ATTTGTTAGT	CCTGCAAAAG	CAGTCTGGTC	CCCAGGCAGG	AAAGGGGTTT	CTTTCTCDDDD
85381	GGGCTGTTAT	TGTTTTTGTT	TAAAAGCAAA	AGTATAAACT	AAGCTCCTCC	CAAAGTTAGT
85441	TAATCCCAAA	CTCAGGAATG	AAAAGGACAG	CTTGGAGGTT	AGACGTTAGA	TGGAGTCGGT
85501	TAGGTAAGAT	CTCTTTCACT	GTAATAATTT	TCTCAGTTAT	GATTTTTGCA	AAGGCAGTTT
85561	CACTGTCCAC	TTCACCTCAC	ATCAGGCCTC	TGACTAGAGG	ATTCCAACAA	TACTTAGGCC
85621	AGGACACCAC	CATGTCTCCT	TATCCACCCT	GAGGGATTCC	AATTTCTCAA	ACAAACGAAA
85681	CTATATATGA	TAGTATGAAA	CTATATATGA	GAAGGAAATT	ATATATGATA	<b>ልጥሮል ልጥጥጥል</b>
85741	GGGTTATCTT	ATTGATTAGA	AGATATTAAA	GTGTGACACT	GCCTGGCAAT	GATATCTCCT
85801	GGTAGTAAGA	ATTTGGCGAA	TTTAGTGAAA	TTCCTGAGGC	TGAACCTCCA	ርጥጥርጥር ጥ አ አ
85861	ATGGAGACAG	TGAGATAATT	TGCCTTACAA	TGCTGAAGTA	AGAATTTTAC	ACAATAATTC
85921	AGACCAACCA	CTTCATGTGG	TACTTGGCCC	GTGGAAGACT	ATCAATGACA	<b>CTT 3 CTTT 3 T</b>
85981	AGTTTATACT	ATTAATGAAT	CCTTTGTTTC	ATTGTTATTT	CCTTCTACAC	GTTGGCCTCT
86041	CIAAAAGAAG	GTAATATTCA	ATACAAATAA	AGTTAAAACA	GCTTGCAGAG	TTGTCCCAGG
86101	GAACTCACTT	AACCACTGAA	GTGTTCAAAT	TGCTTAAGGT	TGACTTTATA	TTCTCCTCAC
86161	TAACCTTTCT	CCTTCTGGTA	TTTCTTCTGA	GAACAGCACC	ACCATCCAAA	GCATCATGCA
86221	AACAGTGGTC	ATCCCAGACC	AGTAATTCTC	AACTCACAGG	GTGCTCCTGC	AGAGATGTAT
86281	TIGAATAGAG	TGGTAGGATG	CTGAAGAAGG	CCACGTAAAA	TTTGGCCAGT	GATCTCCCCC
86341	AGATTTATCC	TGAAGCTAAT	GAAACACAAG	TGTAAGGGCC	TGTACTTCCA	AGGTGCAGAG
86401	AGGGGCCCTA	CAAATGTGTT	AGTTTGTCTC	TCTCTCTCTC	TCTGATTTTA	AAATTTCCAG
86461	TATTAAGGTA	CTTTAATCAC	GGATGGTTCA	GGCTGCTATT	TTCACTCAAT	
86521	ATTAAAATCA	CCATTGTCTG	ATTATGTTAG	AATCCTGATG	AAAATATTTG	GAATTTCACT
86581	AAGAGAAAGT	TTAGTTGAAG	ATGTATCTAG	TATGGGGATA	ATAAGTTACG	TCATTTCCAT
86641	AIGIGATCAT	GTGTACTTCA	TTCGTTGCCA	GCCAATCTGA	CCTAACAATC	COTTONNOCA
86701	GGCCGGGCGC	GGTGGCTCAC	GCCTGTAATC	CTAGCACTTT	GGGAGGCCGA	GACGGGGGGA
86761 86821	TCACGAGGTC	AGGAGATCGA	GACCATCTTG	GCTAACACGG	TGAAACCCCC	でですべて ひっと カ
_	AAATACAAAA	AATTAGCCGG	GCGTGTTGGC	GGGCGCCTGT	AGTCCCAGCT	ACTTGGGAGG
86881	CIGAGGCAGG	AGAATGGCAT	GAACCTGGGA	GGCGGAGCTT	GCAGTGAGCC	GAGATCGCGC
86941	CACTGCACTC	CAACCTGGGA	GACACAGCGA	GACTCCGTCT	CAAAAAAAAA	222222222
87001	IGGCTTCAAG	GAATGTTCCT	ACTGCTCACT	GGAATAACTC	<b>Δ</b> ССТ <b>Δ Δ Δ</b> ΤΤC	CTCCCAACAT
87061	GCAGGTCTAG	ATAAAATGTT	ATGACATCTA	AGTATTCAAA	ACACATTCCC	AGCACTGAGA
87121 87181	GIGAGIGICI	AGTGGAGAGT	AGAAACGTAT	AGAGCCAGAA	GCTAGTCTGG	AAACAATTCT
87181 87241	TACAAAGTTT	ACAACTTACA	TGTGAAAGGA	GCTTAACAGA	GGATTTTCCA	AATTTGAAAA
87241 87301	CAATCCTAAA	AACTTACTTG	ACATTACCAA	TAATGTGTTT	TGAAACTGAA	<b>ልጥልርጥጥርጥል አ</b>
87361	GTTATGAAGA	AAACATATTA	TCATCAGCCA	CCCTGGAGGA	AAGATTGAAT	サーマン サーマー とり
87421	TIACCTATAG	ACAACATTAC	AAAATAATTT	CGATCTGAAG	ATGGAATCAG	AGTATTCAGT
0/421	CAAAACTACA	GGAAAATATA	CTTGGTAGTG	TCATATTCAG	AAGTTAATAA .	AATATGCTAT

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87481	TTTCTGAATT	TTGTGATGG	TGTTGTTTTG	TCAGCTTTTA	TAAAATTGGA	ATTTGATTTT
87541	ATTTTCCCAT	TATAAATTTA	TATTTACAGT	CTGCAGTACT	TTTCCATTT	TAATTTTACA
87601	TTATAGCTTT	TAATAGTTAA	CAAGTTGTAA	AAGGTTTGAT	CCCCAGAAAA	CCTTGATCTA
87661	CCCCCTCAGT	TAAGTATACT	AATATATTTA	GAAAATGGAT	GAAATCAGCA	TTTGAATATT
87721	TTTAAATATI	TATTAAAAGA	GGACATGGGT	AAAAGAGCTT	TGCAGTTGCC	ACCCTTCATT
87781	CTCAAATTCC	CTGGATAAGG	ATGACCGCAT	AATCTTTGGA	TGGTCATACC	CAAGTCTTGT
87841	GTATTTGTTA	CATAAATCTA	TTTAGTGGAC	TTTTGGCAGT	GTGTACTGAG	GCCAGTTTCT
87901	TCCACCTGAG	CTCTGACTCC	ACCTCCAGCA	GCCCAAAACC	AATACTGAAT	TTTGGGGTCA
87961	GCTATTGTTT	TTGTGGACTT	' AGGTAACTAC	ACACACATTG	ТСТТТАТСАТ	AGCTTTAATA
88021	ATACTGCCAT	CAGAACTAAA	ATTGTCACGT	GGATTAAAAG	GAGTGACGGT	GGTGTCCCCA
88081	GGAGCCTTTC	AATATGTAAG	TATTTACACA	TATACATGCT	AAAAAGACCC	CTAGGAATTT
88141	TTTTAACAAG	GGCAAAACAG	TAACTCAGCT	TGTTTTCTCG	CAGTAAAACC	GGTTGAAAAG
88201	GCCTGATAGA	CTTGTCTGCA	GTTACAAAAC	TTGTGTGTAG	TTATCACCTT	TATATCTCCT
88261	GGAAACTAAC	ATAGACAACC	GAATGGGTTA	CAACTGTTTT	TAAGTGAAAT	TGTGAGTGGG
88321	TCTGAAAAGA	GCCTTTTCAA	TGAGGAAGAA	ACGGGCAGAC	TTATGCCCTT	TCCCCACGA
88381	TGCGACGTGC	CAGCTGGATA	TCTTTGGGCA	TGATGGTGAC	GCGTTTAGCG	TGARTACCCC
88441	ACAGATTGGT	GTCTTCGAAG	AGTCCCACCA	GGTAGGCCTC	GCAAGCCTCC	TGCAGCGCCA
88501	TCACCGCAGA	GCTCTGGAAA	CGCAGGTCGG	TTTTGAAGTC	CTGGGCGATT	TCTCGCACCA
88561	GGCGCTGGAA	CGGCAGCTTC	CGGATCAGCA	GCTCGGTGGA	CTTCTGGTAG	CGACGGATTTT
88621	CGCGCAAGGC	CACGGTGCCC	GGGCGGTAGC	GATGAGGTTT	CTTCACGCCA	CCGCTCCCC
88681	GAGCGCTCTT	ACGGGCTGCT	TTAGTAGCAA	GCTGCTTGCG	CGGAGCTTTC	CCGCCCCTTAC
88741	ACTTGCGAGC	TGTTTGCTTC	GTACGAGCCA	TTTGCAATGA	GAGCACACAC	AAAAGTGTAG
88801	TGAACTGAGA	GCAAGTGGCC	TTTAAATATA	GTGAGAAACA	TTCTGATTGG	TCCTGTAATA
88861	TTTCAAAAGT	CCCGCGCGAT	AAAATCATTG	GCTGAAGAGT	GACCAGACTG	ATTGGTTCAT
88921	TACTAGACAA	TCTTATTGGA	TGAGTTGCCC	CACCGCCCAT	CCTGTCCTTT	TCGTTTCAGT
88981	TATCTGCAGC	GACAAATTGT	CTAAAATTCT	AGTTCATCCA	GTCCCAAAGA	ACAGAGTGTA
89041	TAACAAGGTA	TCTAAGGATT	TTTAAAATGT	AAATTCCGAT	TCAGTAAGTT	TGAGTGGGAC
89101	TTGAAATTCT	GCATTCCTGA	CAGTCTCGCA	AGTTATCAAT	GCTGGTGAAC	ACTCACTAAA
89161	CCACCAGAAA	CGTTCAGACT	CATGTCGGGA	AATAACGCTT	ATATTCAGAG	AATGAGATTC
89221	CATGCTATTT	TGTTACTGGC	GAACAGCAAG	TTTCCTTGCC	CTTTGTTTTC	TAAGTCCAAG
89281	TCACATTCCC	ACCCTGCCTG	TTCTCAAAAT	GTCTTATTTT	GGTTGGCCTT	AAGTTTCACT
89341	TTGTATACTC	TAAAATGTAC	TTTCTAAAGG	AAGGTGTTAT	TTTCTCGAAA	CTTAACTTTT
89401	TAACACCATT	AGGCTAGGGG	GGCGGTGGCT	CACGCCTGTA	ATCCCAGCAT	TTTGGGAGGG
89461	CGAGATGGGA	CGATCACTAG	AGGCCAGGAG	TTCAAGACAA	CCCTGGCTAA	AATGGTGAAA
89521	CCCCGTCTCG	CATAAAAATA	CAAAAACTAG	CTGGGCGCGG	TAGCAGACGC	CTGTAATCCC
89581	AAGTACACAG	GAGGCTGTGG	CATGAGAACC	GCGTGAAGCG	GCGGGGTGGA	GGTTGCAGTA
89641	AGCCGATATC	GCGCCGCTGC	ACTCCAGCCT	GGGTGACAGA	GCTAGACTGT	CTCAAAACAA
89701	ACCAATCCAA	ACGAAAAGCA	AAAAATACCC	TAACAGAAGC	AAGTTATCAT	CCTTTCTTGT
89761	GTAACTATGG	ACGGCTCTGA	AAAATGCCGT	TTCAAGTGTA	AGCTACGTTT	TCTGATTTGA
89821	GTGTTTACTT	GACCTTGGCC	TTATCGTGGC	TCTGTTATTT	TGGCAACAGG	ACGGCCTGAA
89881	TATTGGACAG	GACGCCTCCC	TGAGCAATAG	TGACGTTGCC	CAGCTGCTTG	TTGACCTCCT
89941	CGTCGTTTCG	GATGGCCAGC	TGCAGGTGGC	GGGGGATGAT	GCTGCGGGTC	TTGTCACGTA
90001	TGGCGCTGCC	CACCAGTTCT	AAGATCTCGG	CGGCCAGGTA	CTGTAAGTAC	ACTGGCGCAC
90061	CGGCTCCGAC	CGGCTCAAAA	TAATTGCCCT	TTCGAAAAAG	ATGACGGACT	CTGCCCTATT
90121	GGGAACTGCA	AGCCCGGTAG	CGACGAACAA	GTTTTTGCTT	TAGCTCCATT	TTCCACGTCC
90181	GCAAATAGCG	ACCTATGAAA	GCAGCGGAAA	ACTGTGAAAG	ACAAGCAAGC	TGGAATGGCG
90241	CCTGAACAAA	TCCTTTTATA	CAAACTGCAA	GGCTGCAATA	GGAAGCTATC	CTATTGGTCA
90301	ATTATGTTTG	GTGCTTTATC	CAATAGAAAA	AGATAACATA	AATTCCATAT	TTGCATAAAC
90361	CCCACCCCTC	AGTGAAACCG	TGTTTCTTTT	GTCCAATCAG	AAGTGAGGAA	TCTTAAACCG
90421	TCATTTGAAT	CTCAGGACTA	TAAATACATG	GGCTCTGAAC	TGTTCTCTGT	ACTACTCTGT
90481	AGTGGAGAGT	GTTAGTAGCT	TTTCTATTCT	GTTTAGGAAT	AGCAATGCCT	GAACCCTCTA
90541	AGTCTGCTCC	AGCCCCTAAA	AAGGGTTCTA	AGAAGGCTAT	CACTAAGGCG	CAGAAGAAGG
90601 90661	ATGGTAAGAA	GCGTAAGCGC	AGCCGCAAGG	AGAGCTATTC	TATCTATGTG	TACAAGGTTC
90661	IGAAGCAGGT	CCACCCCGAC	ACCGGCATCT	CATCCAAGGC	CATGGGGATC	ATGAATTCCT

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90721	TCGTCAACG	A CATCTTCGA	G CGCATCGCG	G GCGAGGCTT	TCGCCTGGC	CACTACAATA
90781	AGCGCTCGA	C CATCACCTC	C AGGGAGATT	C AGACGGCTG	r GCGCCTGCTC	CTGCCTGGGG
90841	AGCTGGCTA	A GCATGCTGT	G TCCGAGGGC	A CTAAGGCAG	ר דבררבבנידבנ	י אכידאכיסיימייא
90901	AATAAGTGC	T TATGTAAGC	A CTTCCAAAC	CAAAGGCTC	TTTCAGAGCC	ב ארכידא כידידיתים
90961	ICACAAGGA	G AGCTATAAC	C ACAATTTCT	F AAGGTGGTG	TGCTGCTATA	CTCTTTCACT
91021	I CIAGAGGA	r caactggaa'	r gttagcgaa	ACAAGTTTT	GAGCCAAGGT	* TARCTTCCAC
91081	GGGGCCGTG	C GCGGTGCCT	C TTGCCTTTA	A TCCCGGCAA1	TTGGGAGGC	GAGGGGGGGG
91141	GATCACTTG	A GGTCGGGAG	r TCGAGACTA(	CCCGGCCAAC	TATGGCGAAAC	CCCCCCCCCC
91201	CIAAAATAC	A AATGATAGA	GGTCGTGAT	GCGCTCTTTC	TCATCTGTCT	TACCARROTT
91261	CITICITOC	CCTGGGTAA	CCTTCGGGT	L CTATGTATAL	א בארתייתים איז	NACCTON CON
91321	CICCLCCC	I GGTCTAGTA	C AGGAAACTT(	CCTTTCTGG	TAATCAACCA	CCTAATCCAA
91381	I TCAGGGTA:	r AGTGTTCCT(	F TGGGGGTCAT	TAGCCGTTAR	\ \CTTCTTCTC3	CATCCCCCCC
91441	AGGGGAGCA	S AAAAGTCTAI	A GCGACAAAAC	GGCATGTAGG	GATATTTCCT	CCTCCACCOON
91501	GCCTATGCT	TAAATTCTT	CTTCAAGTAT	TGAGGAAACA	ATAAGCGAAG	TCTGATTTCC
91561	CGGGCGCCTT	TATACGGAAT	ATTTCCCGCT	CCACAAAATG	ADDTCCCACT	AGTTTTGAGT
91621	TATAATTGTT	TATCAATGAC	AACAGCTATO	TAGTTTACAT	, שהבהנט הכרעם! המהנה המהנה	TCCCAGAAAT
91681	CCAGATTCC	ATTTCCTAAC	CCACTTAACC	TTCTCATTCTC	CACCTCTCCC	AGATACAAAA
91741	GGGTTTGGAT	TTTGTGCCCT	TCCCCATCTC	GCGCCACTCC	. DARCOTTIGEG	AGGAGGGCCC
91801	CACTTGGAGA	GGGAAATCTT	TTTCGAGAAG	TCCAGGACGC	CARRACTIACT	ATAGCTAAAA
91861	АААААААА	AAAAAAGGCA	GGAAGAGCAC	TAGTTGAGGA	GGAGGACTCA	ATGGGCCAAT
91921	TCTGGGGCTG	GGGCTGGGG	AAGAAATGCA	AGAAGAAAAG	DCAGGACICA	ACTGCACAGT
91981	AAGCAGGAGG	GGGTGGGGGA	ATCGGAGGG	AGTATTTTCA	CCCD ATTOLIG	GGGCATTATA
92041	TGTAGGTGAC	ATACAGCAGT	GTCTTTGGAT	GAAGAAATAA	ACTITION	ACAGTTCTTG
92101	TTTTTGTTT	GAGAAAGGGC	CTTTCTCTGT	CGGCCAGGCG	CCATCATACC	TCACTGCAAC
92161	CTCGACTTCC	CCAGCTCAAG	CGATCCTCTT	ACTTCAGCCC	CTTGACTCCC	TGGGACTAGA
92221	GAAATGCACC	ACCATACCCA	GTTAATTTTT	TAATTTTTC	TGGACCCAAA	GGGTCTTACT
92281	TTGTTGCCCA	GGCTGGTCAA	GCGAACTCCT	GGGCTCAAAT	GATCCTCCCG	GGGTCTTACT
92341	CCAAAGTCCT	GGGATTATAG	GAATGAGTCA	CCGCGCCCGG	CCCACATTTA	ATTTTTAAGA
92401	ATCTTTTAAA	AGAGGTTCTG	GGCCGGGTGT	GGTGCAGCTC	ACGCCTGTAA	ATTTTTAAGA
92461	TTGGGAGGCC	AAGGTGGGAG	GATCACTTGA	GCCCAGGAGC	TCAAGACCAG	TACCAGCATT
92521	TTAGTGAGAC	CTTTTGTCTC	CACCAAAAAT	TTAAAAAATT	AACCAGGCCT	CCTCCCAAC
92581	TTCTGTAGTC	CCAAGTACTG	GGGAGGCTGA	AGTGGGAGGA	TCATTTGAGC	CTCCARCCTC
92641	GAGGTTGCAG	TAAGCTGTGA	CGGCACAACT	GCACTCCAGT	CTGGGTGAGG	ACACACCOTTC
92701	TCTCAAAAAT	AAAAAATAAA	AAAAAATCTG	GATGCCACAC	AAAATGTCAG	TCAACACCCTG
92761	TAAGTGAAGC	ACTTCCCATC	CTAGTACTGT	ATATGCAAAC	TGCCGTTGTG	1 GAACAACIG
92821	TTGGCTTAAA	AATCTACATT	CTTTTTTTAA	TTATAAAACT	ACCACATCCC	AAAGTGACGC
92881	TACTAAGGAA	TTGAGGCTGC	AGTTTAAGAA	GCTGATATTT	AGGATCTATC	TCCCCAAAAACAT
92941	TGAGACCTGG	TAATATAAGC	ATTTTCAAAA	TGAACTTTTG	GGCCAGGTGA	COTTOTTOTTOTTO
93001	GCCTGTAATC	CCAGCACTTT	GGGAGACCTA	GTCAGGCAGA	TCACTTGAGC	TCACAATT
93061	AGACCAGCCT	GAGCAACATG	GCGAAATCCA	GTCTCTACAA	AAAATTAGCA	CCCCCTCCTC
93121	GCATATGCCT	ATAGTTCCAG	CTACTATAGA	GGCTGAGGTG	GGAGGATTAC	TTC3 CCCCC
93181	GAGGCAGAGG	TIGCAGCAAG	CCAAGATCGC	GCCGCCACAG	CCTCACCCAC	707777777
93241	ATGCACCCAC	GCCCTAAAAA	AAAGCATGAC	TCATTAAAAA	AAAAAAATTT	ACCCCCTCCC
93301	GGTGGCTCAC	GCCTGTAATC	CCAGCACTTT	GGGAGGCCGA	GGCGGGCGGA	TCACCACCTCGC
93361	AGGAGATGGA	GACCATCCTG	CTTAACACGA	TGAAACCCCG	TCTCTACTAA	AAATACAAAA
93421	TAATTAGCTG	GGCGTGATGG	TGGGCGCCTG	TAGTCCCAGC	TACTCGGGAG	CCTCTCCCCAAAA
93481	GAGAATGGCG	TGAACGCGGG	AGGCGGAGCT	TGCAGTGAGC	CGAGATCGCG	CCACCCCAC
93541	CCAGCCTGGG	TGACAGAGCG	AGACTCCGTC	TCAAAAAAA	ασασασασα	AAAATTAAA
93601	AAATATGAAG	TTTTGAAGCA	GAAATTATTT	TGTCGTATGT	ממידעררם יום מ	<u> Հահահահան</u> ան
93661	TGCCTGCCTT	CTTCCTTTGT	TACAGAACTC	CAACACTTAC	CCAAAGGTAG	CTCTTCCCTC
93721	AGGGITTCTG	TACTATAGTC	CCTTCTGTGG	TGGCCAGAAA	TATGTTACAG	CAAACACCTC
93781	CCCATCCAGA	CCCCAAGAGA	GGGTTCTTGG	ATCCCGCGCA	AGAAAGAGTT	CARAGAGGIC CARAGAGACA
93841	CCGCAGTGCA	AAGTAAATGC	AAGTTTACTA	AGAAAGTAAA	GTGGTGDDDC	CACAACTACT
93901	CCATAGACAG	AGCAGGACAT	TCCCGAAAGT	AAGAGGAGGA	AGGCATCCAC	COUNCOUNTS
				ADDADGAGGA	MOGCAT CCAC	CCTAGGTACA

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93961	ATACTTGTA'	T ATATGGGGA	G ATGTGCTCT	G CTACAAGTT	T GTGATAAAG	ATTAATTTTC
94021	TTAGTTACT	A TATTTTGCA	A GAATCAACA	TATTATCTT	ייינגממטמממ ד	ATTACT
94081	TTGTTCTCC	A GATATAGGG	A TATCTGGAC	A CTCCTAAGT	TGAGTCTGT	TAGTAAACAT
94141	TATTTATTT	G TTCCCTTAA	CCGTAAACAT	TAGAAGCTAG	GAATGACTC	CTTTCTGGGA
94201	ATGCAGCCC	A GAAAGTCTC	A GCCTCATTT	CCTAGCCCT	ב סוטוניטאניטאני	GAGTTACTCT
94261	GGTTCAAGT	A ACTCTGACA	C TTTTCTTCTC	TTTTTTTCT		TCCTTTATTT
94321	TTTATTTTT:	r atttttgaaj	A TAAGAAATCA	AGAATACTT	ATGTTTCATC	TAAAACAATA
94381	CCCATAATT	3 ATAAGCCAA	A ACAAAAACCI	AGGTCTTCT	ACTCDABACT	1 DCCDTCTTTTT
94441	GCTGTCTCTC	3 CTGATACTC	GCTGATCGTT	AATAGGTAA	ממסממסמד ז	CCCTTCCTT
94501	GICCCCCTC	4 GITTATTAC	: ATTAGATCA1	T ATGCCTACTO	דמידמיודמייי ל	TARTCORORS
94561	CIATGUATT	CACAAAACT	r gccataaaa	TTCACAGGTT	TCCCGCTTCC	
94621	CATTTCCGA	A GGGTCCCATO	IAATATAAAI	CTTATATTAZ	ATACATTCT	אירוייייייייייייייייייייייייייייייייייי
94681	CTTGCTAAT	TTTTTTTTT	TTTTTTGAGA	CTGAGCCTTC	CTCTGTCACC	CAGGCTGGAG
94741	TGCAATGGC	G CGATCTCGG	TCACTGCAAC	CTCCGCTTCC	CAGGTTCAAG	CGATTCTACT
94801	GCCTCGCCCT	CCCGAGTAGO	TGGGACCACA	GATACGTGCC	ACCATGCCCC	GCTAATTTTT
94861	GTATTTTAG	TAGAGACAGO	GTTTCACCGT	GTTGGCCAGG	ATGTTCTCAA	TCTCCTTACC
94921	TCGTGATCCG	CCCGCCTCGT	CCTGCCAAAG	TGCTCGGATT	ACAGACGTGA	GCCACTGCAC
94981	CCGACCAATO	TGTCTTTTT	TAGAGGGGCC	TCAAGCATGA	ACTTACTGAT	GGGTGAGAAA
95041	AACAGAATTT	TCTTTTCCCC	TACAATATAA	ACATTAATTO	TAATGTTATC	ATTCAGGACA
95101	TTTTGGTGAC	CAATCTTACA	GAAATTTTAT	CTTGTGCAAG	TCTATGCAAA	CCAATATCTA
95161	AATCTTCTAT	AAGTGAGATT	GTATTTCACT	TTTCTAGTAT	CCTTTTAAAT	TANTANANCA
95221	GATTCTAATG	ATTATTTTCA	TTACTGCATT	TCATTGTAGG	GAAGTAGATA	ATTCCCCTTT
95281	ATTCACTGAC	CTTCGCTTTT	' TAAAAATTTA	AACCATGTTA	CCATGAAAAT	COTTTTCACT
95341	ATTTCTCTAC	ACACAAGATI	' GCTGTAAGGG	CAAAAATAGA	GATAGGAATC	ATGCATCCAT
95401	TGATATACAT	ATTTTGATTT	' TTAATACATG	TTACCAAGTT	GCCTCCTGAA	GGTCTGTTTA
95461	CACTCTCACC	AACAGGGTGT	TTTTTCCTGA	CTTCCACAAA	TGCTCTTGAA	CAGTGGGTGT
95521	GTTAGTCTGT	TCAAATTGCC	GACATGAACA	ATTAAATCTC	ATTGTTGTTT	ע ע ע אוויידיידיע ע ער ע
95581	GACAATTATT	GTTTGAGACT	GCACATTTTG	ATAATAACAT	TTCTTCTATT	ATGGTTTGAT
95641	TACTCATGAT	TCTTGCCCAT	TTTCTTTTGG	GATGTTGCCT	TATGTACATT	מידית ממידידים מ
95701	GATAGCTCCA	TGTATTAAAA	GATTATTAAG	TTTGAGGGCT	TATGATATGT	Састтасатт
95761	TCTAAGATTT	TITTTTT	TTTTTTGAGA	CGGAGTTTCA	CACTTGTTGC	CCAGGCTGGA
95821	GIGCAATGGT	GCGATCTCGG	CTCACCGCAA	CCTCCGCCTC	CAGGGTTCAA	GCAATTCTCC
95881	TGCCTCAGCC	TCCCCAGTAA	TTGGGACTAC	TGGCAAGCGC	CACCACGCCT	GGCTAATTTT
95941	GTATTTTTAT	TAGAGATGAG	GTTTCTCCAT	GTTGGTCAGA	CTGGTCTCGA	ACTGCCGACC
96001	TTGGCTTAAA	AATCTACATT	CTTTTTTTAA	TTATAAAACT	ACCACATCCC	ССАВАВАСАТ
96061	TACTAAGGAA	TTGAGGCTGC	AGTTTAAGAA	GCTGATATTT	AGGATCTATC	TCCGGAGAAG
96121	TGAGACCTGG	TAATATAAGC	ATTTTCAAAA	TGAACTTTTG	GGCCAGGTGA	GGTGTGTCAT
96181 96241	GCCTGTAATC	CCAGCACTTT	GGGAGACCTA	GTCAGGCAGA	TCACTTGAGC	TCACAATTCC
96301	AGACCAGCCT	GAGCAACATG	GCGAAATCCA	GTCTCTACAA	AAAATTAGCA	GGGCGTGGTG
96361	GCATATGCCT	ATAGTTCCAG	CTACTATAGA	GGCTGAGGTG	GGAGGATTAC	TTGAGCCCGG
96421	GAGGCAGAGG	TTGCAGCAAG	CCAAGATCGC	GCCGCCACAG	CCTGAGCGAC	NCA BECBCSE
96481	AIGCACCCAC	GCCCTAAAAA	AAAGCATGAC	TCATTADADA	ידידיגגגגגגג	ACCCCCTCCC
96541	GGIGGCICAC	GCCTGTAATC	CCAGCACTTT	GGGAGGCCGA	GGCGGGCGGA	TCACGAGGTC
96601	TARTACCEC	GACCATCCTG	CTTAACACGA	TGAAACCCCG	TCTCTACTAA	AAATACAAAA
96661	GAGAATCCCC	GGCGTGATGG	TGGGCGCCTG	TAGTCCCAGC	TACTCGGGAG	GCTGAGGCAG
96721	CCACCCCCC	TGAACGCGGG	AGGCGGAGCT	TGCAGTGAGC	CGAGATCGCG	CCACGGCACT
96781	AAATATGAAA	TTTTCRRCCG	AGACTCCGTC	TCAAAAAAA	AAAAAAAAA	AAAATTAAAA
96841	TGCCTGCC	CTTCCTTTCC	GAAATTATTT	TGTCGTATGT	TCTTTCATAA	ATTTTTTGCC
96901	AGGGTTTOTO	TACTATACT	TACAGAACTC	CAACACTTAC	CCAAAGGTAG	CTGTTGGGTC
96961	CCCATCCACA	CCCCARGEC	CCTTCTGTGG	TGGCCAGAAA	TATGTTACAG	GAAAGAGGTC
97021	CCGCAGTGCA	DAGTANATO	AACTTCTTGG	ATCCCGCGCA	AGAAAGAGTT	CAGGGTGAGT
97081	CCATAGACAG	AGCAGGACAM	TCCCCAAACT	AGAAAGTAAA	GTGGTGAAAC	GACAACTACT
97141	ATACTTGTAT	ATATGGGGAG	ATCTCCTCCTC	AAGAGGAGGA	AGGCATCCAC	CCTAGGTACA
		n. coconc	AIGIGCTCTG	CTACAAGTTT	GTGATAAAGG .	ATTAATTTTC

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97201				TATTATCTTT		
97261				CTCCTAAGTC		
97321				TAGAAGCTAG		
97381				CCTAGCCCTC		
97441				TTTTTTTCTT		
97501				AGAATACTTG		
97561				AGGTCTTCTA		
97621				AATAGGTAAT		
97681				ATGCCTACTG		
97741				TTCACAGGTT		
97801				CTTATATTAA		
97861				CTGAGCCTTG		
97921				CTCCGCTTCC		
97981				GATACGTGCC		
98041				GTTGGCCAGG		
98101	TCGTGATCCG	CCCGCCTCGT	CCTGCCAAAG	TGCTCGGATT	ACAGACGTGA	GCCACTGCAC
98161	CCGACCAATC	TGTCTTTTTG	TAGAGGGGCC	TCAAGCATGA	ACTTACTGAT	GGGTGAGAAA
98221	AACAGAATTT	TCTTTTCCCC	TACAATATAA	ACATTAATTG	TAATGTTATC	ATTCAGGACA
98281	TTTTGGTGAC	CAATCTTACA	GAAATTTTAT	CTTGTGCAAG	TCTATGCAAA	CCAATATGTA
98341	AATCTTCTAT	AAGTGAGATT	GTATTTCACT	TTTCTAGTAT	CCTTTTAAAT	TAATAAAAGA
98401	GATTCTAATG	ATTATTTTCA	TTACTGCATT	TCATTGTAGG	GAAGTAGATA	ATTGCCCTTT
98461	ATTCACTGAC	CTTCGCTTTT	TAAAAATTTA	AACCATGTTA	CCATGAAAAT	GCTTTTCAGT
98521	ATTTCTCTAC	ACACAAGATT	GCTGTAAGGG	CAAAAATAGA	GATAGGAATC	ATGCATCCAT
98581	TGATATACAT	ATTTTGATTT	TTAATACATG	TTACCAAGTT	GCCTCCTGAA	GGTCTGTTTA
98641	CACTCTCACC	AACAGGGTGT	TTTTTCCTGA	CTTCCACAAA	TGCTCTTGAA	CAGTGGGTGT
98701	GTTAGTCTGT	TCAAATTGCC	GACATGAACA	ATTAAATCTC	ATTGTTGTTT	TTATTTTTAA
98761	GACAATTATT	GTTTGAGACT	GCACATTTTG	ATAATAACAT	TTCTTCTATT	ATGGTTTGAT
98821	TACTCATGAT	TCTTGCCCAT	TTTCTTTTGG	GATGTTGCCT	TATGTACATT	ATTTTAAATA
98881	GATAGCTCCA	TGTATTAAAA	GATTATTAAG	TTTGAGGGCT	TATGATATGT	CAGTTACATT
98941	TCTAAGATTT	TTTTTTTTT	TTTTTTGAGA	CGGAGTTTCA	CACTTGTTGC	CCAGGCTGGA
99001	GTGCAATGGT	GCGATCTCGG	CTCACCGCAA	CCTCCGCCTC	CAGGGTTCAA	GCAATTCTCC
99061	TGCCTCAGCC	TCCCCAGTAA	TTGGGACTAC	TGGCAAGCGC	CACCACGCCT	GGCTAATTTT
99121	GTATTTTTAT	TAGAGATGAG	GTTTCTCCAT	GTTGGTCAGA	CTGGTCTCGA	ACTGCCGACC
99181	TCAGGTGATC	CACCCGCCTC	GGCCTCCCAA	AGTGCTGGGA	TTACAGGTAT	GAGCCACTGG
99241	GCCCGGCCAC	ATTTCTAAAT	TCTTTATAAG	TATAAATTCA	TTCAATCTTC	ACCAAAACTC
99301	AATGAAGTGT	GAGTACTATT	ATTATCATTG	TTTTACAGAT	CAAAACAAGT	AATACAGTCA
99361	CTTACTGAGT	TCTATACACC	TGGTAATTTT	TTTGTTTCGT	TGTTCTATCA	ATTATTGGGG
99421	AAGGGGTGTT	GAAATCTCTA	CCTTTAAATC	ATGTATGTGT	CTATTTCTCC	TTTCGGTTCT
99481	ATCAGGTTTT	GCTACACATA	TTTTGCAGTT	CTGTTATTTG	GTGCATATAC	ATTTAGAATT
99541				ATCATTATGT		
99601	GTAATTTTCT	TTGCTCTGAA	ATATACTTAT	CTGATATATC	ATCCAAAAGA	CCACCAGGAT
99661	GGCTAAAGAG	TAGAAAGGAG	AGATTTACTG	GCAATACTAA	TTTGCAAGCC	AGGAAGAGAT
99721	GGTCCCAGAA	CCTGCCAAAA	TTACTCTCTC	TTTGGGGAGA	AGGAGCAGGT	TGGTTATTTT
99781	TATGCCTCAT	AGGCTATATA	TTACACAATA	GAGTCATACA	TATTTAGCAC	GTTTGGGGGG
99841	ACAGCTATAT	ATATTATGAG	GGGTGCCAAG	TGCATTCACA	ATGGATAAAC	ACGTGTAATA
99901	TACCTCCCAT	GTTCACTTCG	AGGTTAAATT	TTGGTTAAAA	TGAGGTAGAA	TTTAGGTCTT
99961	TACATCACAA	GGTGAACTAT	AGGAACAAAG	TTTACGTGCT	GCCTCTAGCA	GCTGGCTGAA
100021				AATAGAATGT		
100081				AATCAGAGTT		
100141				TTTTTGGTAG		
100201	GCCATGCCAG	CCAAGCCATG	AATGCTCTAC	CAGTAGGTAA	CTTTGTTTGC	TTAATCTTAG
100261	AGTCTGTCTT	AGTTGGTATA	GGGGCATCTA	TTTTGGTCTT	TCAGATCCCA	GATATTATTA
100321				GTTTATATGG		
100381				AAGTGAGATT		

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100441	GTTGTTTTT	T TTTTTTTG	A GATGGAATT	T CACTCTTCT	T GTCCXCCC	G GGGTGCAGTG
100501	GCACAGTCT	C AGCTCACTG	AACCTCCGC	TCCTCICT	C AACCCAGGCIO	CCTGCCTCAG
100561	CCTCTTGAG	C AGCTGGGATT	GCAGCCATG	C GCCACCACA	C CCCCCMARC	r TTTGTATTTT
100621	TAGTAGAGA	C AGGATTCAC	ATGTTGCCC	GGCTGGTCT	C CAACTCOTTAAT	CCTCAAGTGA
100681	TCCGCCAGC	C TCGGCCTAC	AAAGTGCTG	GATTACACC	C GAACTCCTGA	CCTCAAGTGA CGCCCAGCC
100741	AAACTGTTT	T TTTATGGGT	TATTTATAC	D GAILACAGG	T BANGACCTO	GCGCCCAGCC TTGATATCTT
100801	AGGGCTTAA	G TTCATGAAGO	GTAGTGTGG	2 NORCACAII	AATGCAATTA	A TTGATATCTT
100861	GCCAGAAAT	CACTGACAAGG	CAGATTGATT	ANCENIAGIO	L TUTTGGCCCA	CTAAATGTTT ACCTATTGTT
100921	TAACGTGTC'	T ATGTGGGAGO	ATTCAGAATT	C ANIMOGIGA	A AAGGCATTT	ACCTATTGTT AGTTATAGAT
100981	GCTTATATA	CATTTTTAGA	TCACAGAAA	ANTIACCIA	T TAGACTOCAATO	G TAAAACAGGT
101041	TATGGGAGG	C AAAAGAGGTT	TGGCTTGCA	ACCTCCCCT	TAGATTCTGG	TAAAACAGGT TGAAGCCTCC
101101	CTCAGAAAG	A ACAGATGGT	AATGTTTCTT	TTATCATOR	GITAGGTAGG	ACTCTCAGTC
101161	TCTCCTGGA	CTGGGGAAAG	CTATAGAAAA	CTCACCACTT	TAAGTGTCAG	ACTCTCAGTC TAATGGAGAT
101221	TCTCTACAG	TGTAAAATTT	, <u>GININGWW</u>	* ACCORAGE	ATGGCTGCAT	TAATGGAGAT TTCTGCCTGC
101281	TGGCCAAGC	GCAGCCATTT	י רבעכבאוווא	CARGCAGCTT	GCAAGCCCAT	TAAAATATTT
101341	TGATTTCCT	TAGACTGGTG	GCCTTATAA	CAAAGAAATA	TATTTTGGGG	TAAAATATTT CTGACACACA
101401	TACCCTTGCT	CTCTCAACAT	CTTATAINAC	AAAAGGAAGA	GACACCTGAG	ATACTAATTC
101461	CATGCCCTTZ	GCTTCCCAGG	TTCTDCDDCD	. AGIAAGAAGG	CCCTCACCAG	ATACTAATTC CTTTAAAAGT
101521	TAGCCAGTCT	GTGGTATTCT	CTTAGAACA	GIAGGAAATA	AATITCTTTT	TATATTATGA
101581	TCATCTTACE	TGACTGATCC	CTCCTACATO	CACAAAATGO	ACTAAGTAAC	TATATTATGA
101641	TGTTAGAGGT	TCCTCTACCC	DICTIONAL AND	MIACACATAC	ACAGGCCACA	TTTGGAACAT
101701	TTTGAGTATO	ТТСАВТАСТА	TATACAAAIG	1ACTACAAA1	TATATATGTA	TTTTTAAATT
101761	TGTATTCAAT	TICAMIAGIA	ATTACTOR CA	AACTITIGTA	GTCAAAATGT	CATTATAACA
101821	TATGGTTTGG	ATOURIARII	CCTCTANATO	TGTTTTACAT	TCTTTCTTCA	TACTAAGTGA
101881	AGTGAAGCCT	GGTGAAAGGT	TTTTCCAMAIC	TCATGTTGAA	ATGTAATCTC	CAATGTTGGA
101941	TCAGGGTAAT	GGTGAAAGGT	TCACTTOOATC	TGAGGGTGAA	CCCCTCATGA	AGCGCACTCT
102001	GTGACACCTC	CAATGGGTTC	CTCCCTCAGG	TTCACAAGAG	ATCTGGTTCT	TTAAAAGAGT
102061	CACCTTCCAC	CATGATTEGA	ACTUTOCACO	TCTCACCATA	TGATATGCCT	ACTCCCTCTT
102121	ACCTCCTGTA	CATGATTGGA	AGITICCIGA	GGACTTGCCA	GTAGCAGATG	CCTGCACCAC
102181	TCAGTTTCAG	CAGCCTGCAC	ARCCGIGAGC	CAAAAAAAAT	TACTITTCTT	TATAAATTAG
102241	ATATTTACAC	GGATTCCCTT	TOTAL ACTION	AAGAACGAAC	TAACACACTA	AGTCTATTTC
102301	GCTAGTGGGC	AATAGCTCAA	ACCOMMONAC	CCTTTTTCAA	CTTCACAGTA	GCTACTTGTA
102361	TTTTTTTTT	ACTGATTTGG	AGCGTGTTCA	AGGGTGAATT	GTATTATGCA	ATTAACAGAT
102421	GTTGGAGTTG	GTTTTCGCAA	ACCACGAGGC	ATAGATTGTC	TTACTTTCTC	TGCTCCTGGT
102481	CCCCAATATT	TTATTGGGAA	ACAACTTATT	TTCCTCTTAT	ATTTATATGG	AATAAATAAC
102541	AATTTACTGT	TCCCTCCCCA	ATATOTGCCT	TTTGTATGTT	TTTTGAAGGC	AAGTGCCTAG
102601	ACATGCCAGG	TTTTGAAGCA	CITACTGAAA	GGATTGCCAT	CAAGTTGTTT	TGCTAATAGT
102661	TTCTCATCCA	CGCTTGTTGG	CAGRARAGE	TCAAGGTAAC	TTGGATGAGA	AGAAGAGTTT
102721	TTTCTAGTCT	TGGCTCAGTG	ACCERCACE	TACTGATATT	GTGACTGGAT	GTACTCCTGC
102781	ACATATCCAA	GAGTTTTTGA	AGCIACCCTI	AATCTTGGTT	TCAATTTTAT	CTAGCCCTGT
102841	ACTITCTGAA	GGCTCTTTCC	TCA CARDON	ACGATITGTT	TAGGAAGTIA	GAATAGCTGT
102901	AAGTATTTAT	CCACGGTTCC	TCCCTCCCTT	TGGACTTCAA	ACACATCCAG	CATTTTATCG
102961	ACATTCTCCT	CCTTCCTACT	TCCATCCTT	CTTCCTTGCC	TTCAGGTCTG	AATTCAAATG
103021	TATTTTTCTC	GATGAAACTT	TCCATCCTTA	TTTCTATTCT	TTTTTCTTAT	CCCCTTTCTT
103081	GTGCACCTCC	CACAGCACTC	CARCITATE	TCTACATTTT	CATTATGTAT	TTACCTTATT
103141	TGCTATGCTC	CACTACAAGA	CAAGIAGCAC	CGTAAGGAAA	CAGGTTGTCT	GCTTTTTCAC
103201	GCTGAACTAA	CCTGCACCTA	TATACACTOTO	TGGCACTTAG	CAGGTTTTCA	GTAAATATAT
103261	ACATTGATCA	TAATGCTGGA	TATACATCIC	CCTCATGAAC	TCTCTAAATC	CTTCTAATTT
103321	TTGTATGCAG	ATCTTCTTTT	CCMIGIGCIT	TIGIATGATT	TATTGCTCAA	AATCTTTATT
103381	GTATAATCTC	AACGTGCACT TTCAGGGCAC	TATCTCACA	CTTCATGTAC	GTAAGTCCTC	CCTTCTCTGA
103441	TACCTTTTCA	TTCAGGGCAC	CCCACGAGAT	AACTTTTAA	CATCTCCATC	ATGAATCTTG
103501	AGATCATTAT	AAGAAAATGA	ACCCARCEMENT	ACTGATGTTT	ACGGCTATTG	TTGAGGGTGA
103561	TCAGAAGACT	AATTTTGAAA .	CARAGATTG	AATATTGTGA	AGGGAAAGAT	AACACTAGAG
103621	AGTTTCTCTC	TGGGAGAAGG	TANANAACAA Tananamamamamamamamamamamamamamamamamama	ACTAAAAATG	AGCACTTTTA	GTCTCCTGAC
		AATCAAATCC	AIAGITCTGT	GACAGCGTTG	GCTTAGAAGC	AGATTTTTTT

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103681	TTTTTTTTT	TTGAAATGGA	GTTTCGCTCT	TGCCCAGGCT	GGAGTGCAGT	GGCACGATCT
103741	CGGCTCACTG	CAACCTCTGT	CTCCAGGGTT	CAAGCGATTC	TCCTGCTTCA	GCCTATGGAG
103801	TAGCTGGGAT	TACAGGCTCC	CACAACCACG	CCCAGCTAAT	TTTTTGTATT	TTTAGTGAAG
103861	ACTGGGGTTT	CACCATGTTG	GCCAGGCTGG	TTACGAACTC	CTGTTCTCAA	GTGATCTGCC
103921						AGCCAGGAGC
103981	AGATTTTTTT	ACACTCATGT	TTCTTTTTCC	TTCTGTCATC	CTGTTTCAGT	ATAAGCAGAC
104041	CACAGATAGA	AGTAGTAGAT	ACCTCAGAAA	TTCCTGGAAT	AATTAATCCA	CGTTCATCTG
104101	TACTCCATCT	GCTCCTATCT	CATGGAATAT	AAAAGGAAAA	ACACCAAGAT	TTCCCTAGGC
104161	AATCTGTCTT	GATTTTAGGT	TCCTCAACAG	GAGAGCCAGA	CAATGGCTGT	AATAATATTG
104221	TCCCGGCCAA	GGAAAAACTT	CCCCTTTGCC	CTCCCAAGGT	TTATGGAAAA	TTACTGGCAA
104281	AACACAGATT	AACTGGAGAA	<b>AAGGCATATA</b>	TATTTATTTC	ATCACAATTT	TACAGGAGAT
104341	TTTAGAATTA	AGACTGAAAG	ATACAGGGGA	AATTGCCCAT	TTTTATGCTT	AGGTTCAACA
104401	AGATAAACAG	CTGTATAGGG	TACGATCTAA	TGCTAACAGA	CTGAGTGGGG	AAGCCCCGCA
104461	AGGCTTGTCT	GTCAAGATTC	TTCTTGACCT	CTCAGTGCAG	CATTTCTTCC	TTCTGGTTAT
104521	AGGACAAGAC	TCTCTTTTAG	AATGGGGGGT	CTTATGACCT	ACAGGCAAAC	AAGGTAGGTT
104581	AGAGTAATAT	TTTTAGGTTT	TATGGCTGGT	TCTAGGGAAA	AGGAGTTCTG	GTTTGTATGG
104641	CCTACCTTGA	GGAGGAATTC	TGGTTTCTAT	GGCTAGACTT	TGGGGAGAAT	GGGACTTACA
104701	GACAGGAAGG	CAGAAGGTGG	TCAGTGAAAC	ACTTTTATAA	TCATAATCCC	ATTTTGAGTA
104761	TTTCTGTGTT	ATGGAATGTT	TGTTCTCTCA	TTTCCTGAAA	GATTCCAGAG	ACTCCTCATT
104821	CAGTGTTGTG	AAAAAGTTCA	GGAAATGCAA	CTCAAAAATG	TGCCACTTTG	TTACGCTGAT
104881	TTCTTTGAAC	TGAGGGCACC	TAGGAAACAG	TAAATTCAAG	GAAGGGCTTT	CGCTGAACTC
104941	TAATCAAAAA	TTTGAAAATT	AAAAAAAAT	TCAAAAAGGA	ATTTAGTTGT	TAAGATTCAC
105001	TTCCCTGGGG	AATCTCATCA	ACCAGAGAAG	ATTAACTGTA	TCACAGGAGA	GGAGACTGGT
105061		ATCTAAACAG				
105121	TATTTTTCTC	CAAAATCATA	TACTCTCCCC	TAAGTTGCCT	ACATCCCCCT	TCTTTCTCCC
105181	TTATGAATCA	AGAGAGCTTA	TAAGCTTCTA	CAGTTCACTG	GGATTTGGGG	TATTCGCTTT
105241	TCTTCCCTCC	CACTCCCCCT	CCCCTTTTTT	TGTCTTTGAG	ACACAGTCTT	CTGGCTCTGT
105301	CGCCCACGCT	GGAGTGTGGT	GGCTCTATGT	GAACTCACTG	CAACCTCCTC	CTCTCGGGTT
105361	CAAGCGATCC	TCCCACCTCA	GCTTCTCGAG	TAACTGGAAC	TACAGGCGTG	CACTACCAAG
105421	CCCGGCTTTT	TTTTTTTTTTT	TTTCTCCCCC	GTTTCTTTTT	TGGTTATTTT	ACTGGAGACA
105481	GGGTTTCTCC	ATGTTGTCCA	CGCTGGTCTC	GAACGCCTGA	CCCGCCGTCC	TCGGCCTCCC
105541	AAAGTGCTGG	TATTACGGGC	ATGAGCCACT	GCGCCCGATT	TGAAGGACCT	CTTAAATATC
105601		TTGGTCGGAG				
105661	GCACGAGCGG	CTGAAAGTCA	AAATAACCAG	AACAAAACCT	CCACTCATGC	TTAAAAAAGG
105721		AAATCCTAAT				
105781		GGGATTGGGT				
105841	ATGAAGGATG	CAGATTCTGA	TTTCCCATTG	GGTATTTGAC	ATTAGCCAAT	GGGAGAATTC
105901	CTCACAGCCT	ACCTCCAGTC	AGTATAAATA	CTTCTCTGCC	TTGCGTTCTA	ATGTAGTTTC
105961	ATTACATTTT	CTTGTGGCGA	TTTTCCCTTC	TTATCAGAAG	TAGTTATGTC	TGGTCGCGGC
106021	AAACAAGGCG	GTAAAGCTCG	CGCCAAGGCT	AAGACTCGGT	CTTCTCGTGC	AGGTTTGCAG
106081	TTTCCTGTGG	GCCGAGTGCA	CCGCCTGCTC	CGCAAAGGCA	ACTACTCCGA	GCGCGTCGGG
106141	GCTGGCGCGC	CGGTGTATCT	CGCGGCGGTG	CTTGAGTACC	TGACCGCCGA	GATCCTGGAG
106201	CTGGCGGGCA	ATGCGGCCCG	CGACAACAAG	AAGACCCGCA	TCATCCCGCG	CCACCTGCAA
106261	TTGGCCATCC	GCAATGACGA	GGAGCTTAAT	AAACTTTTGG	GGCGTGTGAC	CATCGCGCAG
106321	GGTGGCGTTT	TGCCTAATAT	TCAGGCGGTG	CTGCTGCCTA	AGAAAACTGA	GAGCCATCAT
106381	AAGGCCAAGG	GAAAGTGAAG	AGTTAACGCT	TCATGCACTG	CTGTTTTTCT	GTCAGCAGAC
106441	AAAATCAGCC	TAACAGCAAA	GGCTCTTTTC	AGAGCCACCT	ACGACTTCCA	TTAAATGAGC
106501	TGTTGTGCTT	TGGATTATGC	CGCCCATAAA	GATGTTTTTG	AGGTGTTTTT	AATGGCTTTG
106561	AGTGTGGCAC	TTTTAGTAAT	TTGTCCTGCA	GAAATTAGAT	CCATAGAAAC	CTCAGGAATT
106621	CTAGGTATGT	GGGAGAAGTG	CCATGCAGCA	CAAAACATGT	TTACAGGGGT	GATTCGCGTT
106681	AAGTTTCACA	CACAGCAGTT	ACTACATTTT	AGAGGAAGGA	AATTATACCC	ATGAGTGCAT
106741	TCCTAACTAT	CTTGAATGGA	AGTGTTAAAA	CCCGCATGCC	CCACACAAGT	TTGAATATGT
105801	CATACCATTT	GCTGTAGCAA	TTAATGGCAT	ACACAATTGA	GAGCACACAC	ATTACCACTG
106861	AACATTTGAG	TATGTATTTC	CCAAAATGAG	CTTTTTTCCA	GTTTGGGGAT	GTTTTGCTTT

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100001	COMPRESSE					
106921						TAACAGCTCA
106981		GAACTCGGGC				
107041		GCCGCCACGC				
107101		AAAAATACAA				
107161		TGATTTCTTT				
107221		TTGTAAATTT				
107281		GCAGCTAAGG				
107341		TTAATCTGCA				
107401		TTTCTTGGGT				
107461		CTCCACAGAC				
107521	GTACTAAACA	GACAGCTCGG	AAATCCACCG	GCGGTAAAGC	GCCACGCAAG	CAGCTGGCTA
107581	CCAAGGCTGC	TCGCAAGAGC	GCGCCGGCTA	CCGGCGGCGT	GAAAAAGCCT	CACCGTTACC
107641	GCCCGGGCAC	TGTGGCTCTG	CGCGAGATCC	GCCGCTACCA	AAAGTCGACC	GAGTTGCTGA
107701		GCCGTTCCAG				
107761		GAGCTCTGCG				
107821	GGCTCTTTGA	GGACACAAAC	CTTTGCGCCA	TCCATGCTAA	GCGAGTGACT	ATTATGCCCA
107881		GCTCGCTCGC				
107941		TAAAACCCAA				
108001		TTTTTTTTTTG				
108061	TTAGACTATG	GTCTTAAAGT	TGATTAACAG	AAATAACGGT	TTGGTCAGTC	TTGCAGTGTA
108121		GACCTTATTA				
108181		AGCTTGCTAT				
108241	AAATCTGGTA	AGTAGTTAAC	TGGCGCTTAC	TAGGCATTTT	TGCAAAGCTT	TGAAAAGATT
108301	AGAAAATTGT	GTCTTGCGAG	TTCCAGTGTC	TTCCTCAAAA	TGCTTAGGAA	GATTTTCTCA
108361		TAGTCCCCTA				
108421	ATATATATAT	ATATACTGTT	AAATTCATTT	GGCTGTTAAC	ATTAACCTGA	AATTTATTCT
108481		GTGAGGCAGG				
108541		GTCAGTCTGT				
108601	ATGTGTGGCT	TTGCTTGTAA	ATAGTCTATC	TGGTTGCATT	GCTTTGTCTC	CTCTAGGACT
108661	ATGCACCATG	ACATGCCACA	TTCTTTTTTT	CAGTACTTCT	TGCCTGTAGT	TATTAAAATC
108721		AAGTTTTAAC				
108781		GTACTGGAAG				
108841		TTAAAATTTT				
108901		TAACTTTAGA				
108961		TCACATTCTT				
109021		AGAATGTAAA				
109081		ACACACACAA				
109141		TTTTAATAAG				
109201	TTATGAAAAT	GAATATGTCA	GTTTGTTTTA	TGATTCGTTT	TTCTTGACTC	TTATACAAGC
109261		GGCATAGACA				
109321	ATGGACTTGG	TCTATGCCAA	GGTGACTACT	CACAAGCTCT	GGGCCCAGCT	GAAGGTCAAG
109381		CCAGTTATAG				
109441	TTCTTTATCT	GTAGGAAACA	AATGTGTTGG	AGGTACTGGG	TCTGACGAAT	AGCATAAAAG
109501	AATAAAGTTA	CATTACTGTC	TGAGGATCAG	ATGGACAGGG	GGTGGTAGCT	CAGTCCAGCT
109561	ATTTTCCACT	CCCTCACTTA	CATTCTTTGC	CCCCTCCTCA	ACAGAACAAG	GATTCTGCTG
109621	TAACTCTTCA	TTGACAGTTG	ATATTTAAAA	ATTAACGAAT	GGATGAAATT	CTCATTTGTG
109681	AAAGAAAATT	TATTGAGCAT	TTTGTATTTG	TGAGTAGTGC	AAACATTTTA	ATATTATATT
109741	AAGAATCTAT	TGTTTTGTAT	TAGAGGAGTA	ATTAAGGAGA	GATTGGAGAC	AAAAAGGGGG
109801	TGTTGTTTGC	AGAATATACC	ATCCAAAAAT	AGACCACTGT	GGGATCAGGA	TTCTTTTGAG
109861	CTAAAGGCAC	TTCAAAAACA	GCATTCAAGA	AGGGAATTCT	TCTAAACTTT	TCTTTCTGAA
109921	AACAGGAGAT	AAAAGTTCCA	ATGTGAAAAA	TGCTCTGCTT	GTACCAGGTG	AAAAGACATA
109981	TTCTTCAGCC	CAGAGGCATA	GATGAGATAA	TTCTGCACAA	ACACAGCAGG	GAGTCATAGC
110041	CGAGAGACTT	CTATACACAA	ACAAACCTTG	TTAAAATAAT	CATATATTCC	TTTAATCTCC
110101	TCATATGGTT	TACTTTCCCA	CAATTGCCTC	TCTTTAACTT	AATGTGAAAG	CATTTAGCTT

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110161	TTGCCATTTC	TTTGGGGCTT	CACTTTTTA	TGAGGGTTCT	CCTGTCCCAT	AAAATTTACA
110221	TTAAATACAT	TTGTATGCTT	' TCATTCTGCT	AATCTGTTT	ATGGCAAATG	AATTATCAGG
110281	TCCAGCTGGA	GACCCTAACA	GAGTAGAGGT	AAAATTTTG	CTCCCTACAA	GATAGAGATT
110341	GTGTGCATTA	AATGTTGTTT	GTTCCCAGTT	GTTCAGTTTC	TCAGGCCTCT	GAGCCGAAGC
110401	TAAGCCATCA	TATCCCCTGT	GAACTGCACG	TATGCCTCTA	GATGGCCTGA	AGTAACTGAA
110461	GAAACACAAA	AGAAGTGAAA	ATGCCCTGTT	CCTGCCTTAR	CTGATGACAT	TACCTTGTGA
110521	AATTCCTTCT	CCTGGCTCAT	CCTGACTCAA	AAGCTCCCC	ACTGAGCACC	TTGTGACCCC
110581	CACCCCTGCC	AGCCAGAGAA	CAACCCCCTT	TGACTGTAAT	י דיידיריר איידיאיי זיניזטאטנאניני	CTACCCAAAT
110641	CTTATAAAAC	GGACCCACCC	CATCTCCCTT	CGCTGACTCT	TTTCCGACTAT	AGCCCGCCTG
110701	CACCCAGGTA	GAATAAACAG	CCTTGTTGCT	CACACAAACC	י רייניייייטקטפייי	GTCTCTTCAC
110761	ACGGACGCGC	CTGAAACAGT	TTAACAGGGT	TTTTCCTGCC	CAGTITICATO	AAAGTGATGT
110821	TATGCTGCAG	GCTGAAGTTT	ACAGCTAATG	СТСТТСААСТ	CROICACAAC	TTTTGGTTTG
110881	TTAGATTTGG	GTGAGATGGC	TAAGATTCTC	AGAGAAAGAA	CINNAMICAG	GGGTGCATTT
110941	TTCAGACTTA	AAAATTTAGC	AGTAGCCCTT	GCAGTTTTTC	CARTACAACE	GATTTACGAA
111001	TGTTTTCAGG	AAATTTAAAA	CAACAGTGAG	AAGCGTGTAT	CUVINGWAR	GATTTACGAA
111061	AGACTTGGCT	ATAGGAAAGC	ACGAATGCTG	CTBTTCTNTT	GGAGAGIIGA	ACTACACTCC
111121	AAGGAATATT	TTCGGACAAT	TTTAACATGT	CACATATCAA	A A C CTTA A A C C	AAAGAGAACA
111181	ACACCTTGTA	CGTTATTACA	GGCTGTGATT	TTABARARA	AAGCIAAACG	GAATCTGTCA
111241	TAGTTGCTGC	TAGCAATATA	GTGTTGGGAG	TANANACAC	AAICCITACT	AATACATACA
111301	TATCCCAACT	CTGAGCAGAT	TTTTTTAAGT	ACTA ACATOT	AMAMIGAGAG	TTCAGGACAA
111361	AATATTTATT	TCTTTTCCAC	AGTCTCTTCT	CATCCCTCCT	TCACATTAAAC	CATATTATGT
111421	TCCCCTGAGT	ATCATCATAA	CCCGATTTAC	ACATCOLICGE	I CACATTAGE	TAATTAAAAG
111481	CCCTCTTCTG	AATGAGACAG	TACAGTGTGA	AGAIGAAGGC	ACGGTTGCAA	TGAGCTATCA
111541	AGGGCTCTGG	CTGGACCAGC	אסוטוטוטא	AGGATAGCAA	CCATTARCAC	CCATCCTCTT
111601	TATGCATTTA	TTTAACACAG	GTTTTACGTG	VCVCVCCACCACC	TOTOLTALACAG	GAGAAAGGTA
111661	CCAAAAAAAG	CAGTTAGCTA	CTTATATAAT	CA ATTCCA CA	ATTROTTANGG	TAATGAAAGC
111721	CGCTAAAGCA	AAGGGATTTA	GGCTAGAATA	TATAACTCTC	TACACARACA	TGTAAAAATG
111781	GCTAGTGCAA	GGTTTGTACA	GAATTCTCTT	GCCTCACCC	TCCTATCCTT	CCCAGCAAGG
111841	TTGCTTTTTT	TAAACTACAG	TGAGAACATC	TTTCATATCA	CAATTTCLII	GAGAAGAATG
111901	AAGAAACAGG	TCAGCTTTCA	AGAAAACATA	AGGCCAGAGT	CAMCTOTOTO	COCCECCTOT
111961	TTTAAGTACC	TTTGAATAGT	CAATATGTCT	TCAAGCAGAGT	CARCITICA	CGCCTGCTCT
112021	CACTCCGGCA	TATTAGTGAA	AGCCCTTAAT	ATAAGCCCTT	ATTANANTE	TCACTCCACC
112081	GTATAAATTC	AGATTCAAAT	AGTAGTGTCG	TABACGGGAG	CCANANACTA	A A COCCA MINA
112141	AAAGTGAAAC	TATTGTGTTC	TCCCTCGCAG	TCCTTAGGTC	ACTECCCCTC	CACCCCCCCA
112201	GCAAAAAGTG	AGGCAGCAAC	GCCTCCTTAT	CCTCGCTCCC	COMMENT	CTCNATAACC
112261	TCCGATGTTC	GTGTATAAAT	GCTCGTGGCT	TGCTTTCTTT	TCGCGTACCT	COTTO
112321	GTCAGCTGGT	TAGACATGTC	TGGTCGCGGC	AAAGGCGGTA	AAGGTTTCCC	TARCCARCET
112381	GCTAAGCGTC	ACCGAAAAGT	GCTGCGGGAT	BACATCCAAG	GCATCACCAA	ACCCCCCAMM
112441	CGGCGCCTTG	CTAGGCGTGG	TGGGGTTAAG	CGAATTTCCG	GTTTCATOR	TCACCACACA
112501	CGTGGCGTTC	TCAAGGTGTT	TCTGGAGAAC	GTGATCCGGG	ACCCCCTCAC	CTACAGGAGACI
112561	CACGCCAAGC	GCAAGACTGT	CACTGCCATG	GATGTGGTTT	ACCCCCTCAA	CIACACGGAG
112621	CGCACTCTGT	ACGGCTTCGG	CGGTTAATCT	TTTCGTCAGT	TTTCTTCCAA	TCCCCCTTTTTT
112681	TAGGGCCGCC	CACTCCCTCT	CAGAAAGAGC	TGTGATTGTA	TITCITCCAA	TGGCCCTTTT
112741	TCAATGGCTT	TACTCGGCTA	TTCTGCCTAG	TATGTAGAAC	TATTATAAAC	CAGTTCCCAC
112801	AGACCAGGTT	GTTTGGTCTG	AGTGGCTGCT	AAAGCAGAAA	TCAGCTAAGT	AAACCACCTC
112861	TCCGAGATAA	GTGAGCTATA	AACTTCAATG	CTATAGTTTT	GACATGTCAA	AAACGAGGIC
112921	GTGCAGCGCG	AGTCCGATAA	ATGAGTAGCT	CAGCTTTTTA	GTTTTDDDDD	CCACCTICAC
112981	GTTATTTGTA	CGAGAGCCTA	AGATGCTAGC	TGCCTGGAAC	TGAGTAGGTC	CONGITUIGC
113041	GGTGTCAGGT	CTGTTTTCCC	AGGCGTATCT	GACTTAACGT	COCCODANCE	ACTIVAMMIQ
113101	GCTTCCCTGG	TAACACCTGC	CGTCCTTAAC	CCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	CGGTAGCCCC	ACA A CCCOMM
113161	ACTTCCATTT	CTAGTTGAGC	TTGGCGTCCT	GCTGAGTGAC	GTCACCTCC	AGAAGCCTTT CCTTCTCTCC
113221	AGTAGGACTG	GCGGTTAAAG	CTGCTTTGCT	ATTTTCACTC	CTCAGGCTGC	recitetede.
113281	AAGCAGGCTG	CCTACGCAGT	TCGTAAATTC	CCACTTAGTO	GACTANGCIGG	CDCDCDCDCDCD
113341	TAAATAAGGA	CTCAAATTTC	TTCTGACTCC	GAGGTCCGTG	CUTCIUNGCON	TARCETTIA
	-				CCUGCYGCTA	TAMONTGGAN

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113401						TCAACCCAAT
113461						GAAAGTTTCA
113521						TCAGCCAAAT
113581	ATTCAGCATC	TTTGATTTAG	TAACAAATAT	ATTGATGGCT	ACTTCAGCAA	AAAAAATCAA
113641	CTTTGTTTTC	TGGTTACTTT	GCTAACAAGC	TTCTCCTGAC	AGGAGGATAT	AGTGAATAGG
113701					GATAAAAATG	
113761	CAGCAGATAA	ATAAATGCTG	AGACCAGATG	AGATGGCTAA	AAACTGAAAC	ATAATGTAGT
113821	GCAGCATTGT	TTGTAATAGT	AAATGAGTGG	CAACTGTAAA	GTTTTCATCA	GAAAGGACTA
113881					CAGCCCTACT	
113941	AAAGCTGTAC	TCCACTACAT	ACTCTGGTGT	ACTCTGGCTC	AGTTCTTGGA	CTCCTCTTTT
114001	CTTGGCTAAC	TCAACTGGCC	TCACCACTTA	CATGCTCTGT	GCTCTGTCAA	ATAGTTTGTT
114061	CAACAGAACA	CCACGGCCTA	GCTGTAAGTG	CCACGTTAAC	TTCTAGCAAT	GCCANAGCCT
114121	GTGATAGTGG	CAGCTTCGGG	CTGTTTCTCA	TTCCCGGGAT	GCCTAACCAC	CTCTCCAAAT
114181	TCTATCAGTT	TGCTTCCACC	CACTTCAAGC	TTCAGAACGA	AACATAGAGC	TTAAGAAATA
114241	TAGGCCCGGC	AAGGTGGCTC	ACGCCTGTAA	TCCCGGCACT	TTGGAAAGCT	GAGCCTGGTG
114301	GATCACCTGG	GGTCAGGGGT	TCGAGACCAG	CCTGGCCAAT	ATTGTGAAAC	CCCGTCTCTA
114361	CTAAAAAAAA	AAAAAATTA	GCTGGGCATG	GTTGCGGGCG	ACTGTAATCC	AAGCTACTCG
114421	GGAGGGTGAG	ACAGGAGAAT	AGCTTGAACT	CGGGAGGCAG	AAGTTGCAGT	GAGTTGAGAT
114481	CGCGCTATTA	CACTTAGGCC	TGGGAGACAA	GAGTGAAACT	GTGTCTCTAA	ATAAGTGTTT
114541					CATATACAAC	
114601					AATATGTTTA	
114661					TAGAACTCCA	
114721					TTCTTACAAC	
114781					TIGITITETT	
114841	CTAGGAGCAA	GCTGCCATGG	CGGTTTGTCT	GAATGACCAC	AGTGACCCCA	AACTCCTCTT
114901	TGTTTTCACT	TTTAATCCCC	CTGTCATACA	CTTTTTTCTCT	ATCCAGCATC	AACIGGICII
114961					CCACTGGCTT	
115021					AGATGACCAA	
115081					AGTGCTGAAA	
115141					GTACTCTGGC	
115201					TGTTTCTGGA	
115261					TAGAAGGGTC	
115321					CTCAAATACC	
115381					CTGCTCATTT	
115441					TTCCCCACCT	
115501					ATATTTTATT	
115561					ACTTTGTTTT	
115621					GTTTATGGAG	
115681					TCCACTAAGC	
115741	GCTAACACAA	TCAATTCCCC	ATCTCATTCC	TTCACCTCC	ACTGCCTGAA	CCACCAGAAT
115801	TGCAGTTTCT	CTTTAGAAA	TCTCCCCCAT	ACTOTACCO	TTGCAAATTA	GCAATCAGCG
115861	TCTTTCTTCT	CARCARGAC	TCTGGGGGAT	TENCENCE OF CR	AGAAGGCCCA	AGCAACATTA
115921	GGGTATGCCT	AAGATGAGTA	TCACATGAGIG	CCARMCOMAM	AGAAGGCCCA	AGGTGGTGGT
115981	AGGCCAGGCA	ABACCTARCA	CCACCTACTC	CTCCCCTTAT	GAACATAGCA	ATGCTATGAA
116041	CATATGGGTA	ATCCATATCC	ACAGCIAGIC	GIGGCTIATI	GTTACAACGA	CTATACCTCC
116101	ATTAAAATTT	TOTABOTATE	CEN CCCCN NE	CIACATIGAC	TCTGGAATTC	AGGAAAGGGA
116161	ATAAATATCT	TTAAAATACC	ANCERRACIAN	CACATARA	CLOCCLOCCA	CONTRACATOR
116221	CTCATTTTTG	CTCCTCTCAT	TOTOTON NA CO	ATCCACA ATC	CACCUACCCT	CCATACCAGG
116281	TGTCAGTTTT	ADATTABACCO	ANGORGOGEN	CTTCTCTT	TOOMONNON	AMOGRACIA
116341	AAAACTGTCA	GGTGGAAAAC	ACCICCOCIA	ANDANACE	CAMCCAMCA	ATCCTGGATG
116401	ATCGTGAGGC	TGDADTCACT	<b>してはかれかなしかな</b>	TCCTACCAA	ANCACCATCAC	TGAAGCTAAC
116461	TTGTGAATAT	TTACTCAGI	COLMINACAN	TATTTCCCT	MAGAGCACAA	TGAGAGGCAT
116521	ATCCCTTTTC	CACCAGAI	CTCR & CCTR C	AMOUN COMES	TUAGUTAACC	TGAAGTTCAC
116521	DGGDDCDTCC	TTTAAAACII	TOCOMECTAG	AIGTACTTAA	CIGGAACACA	TAACTGCATC
**0761	AGGAACATCC	LITHMMACTA	IGGCTACAAT	GGCTTGACTG	GACAAACCCC	AGGCTTCCAG

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116641	GTTTAGCACA	GGTGGCCCTT	CACAGACCAA	CATTGCCTAT	GCTACCAACC	TCATGTCCTA
116701	CCACCCTGCT	TGCATCATTT	CTCTCTCTGC	ATATATAAA	ATATATGTGT	ATGTATATAA
116761	TCAGCTTTAT	TGATATTTAA	TATACCACAA	AATTTGCCCA	CTTTAGGTAC	AGTTCAATGA
116821	ATTTTACCGT	GTTTTCTTAG	TTGTACAACC	ATCATCACAA	TTTAATTTCG	GAATATTTCT
116881	ATCACCCAAA	TTTCCATTTC	TGCGTAAAGG	GGGAAAAAA	AAGGTTAACT	GCTGAAGGCC
116941	GCGGTAACAC	TGAAAAAGGT	GCCTTTTCTC	TCTAAAACAG	ATTTTAATCT	CCCCTGAATT
117001	TAGTGTCCTG	GGTATTCCAG	GAGTCTGAAT	AGGGTTTCAA	TTTTCAGGGT	CTTTTTAATA
117061	GAGTAAAACT	GTATTGGTGG	CGATAAATTT	AGTATTGCTC	TCAGTACATG	ATTGAGGGAT
117121	ACTTAAATGT	CTCTGTGATT	TTATTTCATA	ATCGCTAAAA	GATGGTTTTT	TTTTTTCCTA
117181	AAACAGGGTT	TTTGTTTTTT	CTCAATAAGC	TTCTTAGCTT	CCCCTCCGGC	TCCCTGGCTT
117241	GCCTCAGGAA	ATATTAGCTC	ATCAGTTCTG	ATTGGTTGAC	AGCTACGAAT	GGCCCTCATT
117301	GATTGGGCAG	CGCTTCTTTG	TCCCTTGGAA	ACTAATACAA	ATTTTTAACA	CTACTTTTTT
117361	TCCACTCTTT	CTTCAGAGTT	GGAATATCGT	TGCTCCCCTA	CCCATATGTA	GTGAGTGGAG
117421	GGCAAACTTG	GAGTTCCCCT	AATCTTTCCT	TTTTAGGATG	TCAGCTCAGT	ATCATTCATC
117481	TTAATTACAC	ATTGAGCTTC	TTGACTTAAT	GGATACAGCT	CTTCTTTTGT	TTAGTTGGGC
117541	GGCCCTGAAA	AGGGCCTTTG	GTTCAGAAAT	GCAAGCTGTG	GAGAAATCAG	CAACCTTAAC
117601	CGCCAAAGCC	ATAAAGGGTG	CGTCCCTGGC	GCTTAAGCGC	GTAGACCACG	TCCATGGCAG
117661	TGACTGTCTT	GCGCTTGGCG	TGCTCCGTAT	AGGTGACAGC	GTCACGGATC	ACGTTCTCCA
117721	AAAACACCTT	GAGCACCCCG	CGAGTCTCCT	CGTAGATCAG	ACCAGAGATC	CGCTTCACAC
117781	CGCCACGCCG	GGCCAGACGC	CGGATGGCCG	GCTTGGTGAT	GCCCTGGATG	TTGTCACGCA
117841	ACACCTTGCG	GTGGCGCTTG	GCACCCCCT	TACCCAAACC	CTTCCCGCCC	TTACCACGTC
117901	CAGACATGAC	TTCCCAAGAA	GTGAACCAAG	AGCAAGTGAG	AGAATAGGAA	ACCGATCTTT
117961	ATATATCTAC	GTTACCCCTG	CCCCCACCTC	CAGCGGACAC	AGAGACTGAA	AAGCGCGCAG
118021	GCGGGAAATG	TGACGCCTAC	AGTCCGCTCC	TTTAACCCCT	CCTCCAAGCC	CCAGGAAATG
118081		CGATTGGGGG				
118141	ATGGCCTTTA	TTTTCTTAAC	AGAGCTACAG	GCTTTGAGGA	ACTGGGTTAA	GAATTAAATG
118201	TAAACCCATT	CTGACTCCAG	AATTATTTTA	AGTCGAACTT	TTTTTTTAAC	CGAATCTCTC
118261	TGTCGCCCAG	ACTGGAGTAC	ATTAGAGCCA	TCTCGATTCA	CTGAAACCTC	TGCCTCTCAG
118321	GTTCAAGTGT	TTCTCCTGCC	TCAGCCTTCA	GAGTGTACCT	GGGATTACAA	GCGCTCGCCG
118381	TCGCGCCCGG	CGTGTTTTTG	TATTTTTCGT	AGAGACGGGA	TTCGGCCATG	TTGGCCAGGC
118441		CTCCTGATTT				
118501	TACAGGCGTG	AGTCACCGCG	ACCGGCCGAA	ATCGATTGGT	TTTGAAGCCT	TCAGTAGCAT
118561	TAAAACGAAA	AGTGCTCCCA	ATGCATTCCC	TTTTGTCTTA	AATTGGTTTC	TTACAGCTAC
118621		AAGGTGGTGG				
118681	AGTAATCACG	CCCTCTCTCC	GCGGATGCGG	CGGGCGAGCT	GGATGTCCTT	GGGCATGATA
118741	GTGACGCGCT	TGGCGTGGAT	GGCGCACAGG	TTAGTGTCCT	CAAATAGCCC	TACCAAGTAG
118801	GCCTCGCACG	CCTCCTGCAG	AGCCATCACA	GCGGAGCTCT	GGAAACGCAG	GTCTGTTTTA
118861	AAGTCCTGCG	CAATCTCGCG	CACCAGGCGC	TGGAAAGGTA	GTTTACGAAT	AAGCAGTTCA
118921	GTGGACTTCT	GATAACGGCG	GATCTCGCGC	AGAGCCACGG	TGCCCGGCCG	GTAGCGGTGG
118981	GGCTTTTTCA	CGCCGCCGGT	GGCCGGAGCG	CTTTTGCGGG	CTGCCTTAGT	GGCCAACTGT
119041	TTGCGTGGCG	CCTTGCCACC	AGTAGACTTC	CGAGCAGTTT	GCTTAGTGCG	AGCCATGACG
119101	GAAAAACAGC	ACAGCGGAAC	ACCCAACACT	AGCGCAAATA	CGCCCATGAG	CTGCTCTATT
119161 119221	TATAGTGTGT	AAAGTGCAGT	GATTGGATGA	TAGAAGACGC	TAAATATGAC	GTTACACACT
119221	CTGATTGGTC	TATCTTTAAG	CCAGCAACAA	TCGTGCAGTT	TCACCGGCTA	CTATATTCTA
119261	TTCCAACTCT	ACAGATGATT	ATTTAAGTGG	TATTTTATTA	CTACTATTAT	TTTATTTTAC
	ATTOCACTOR	TTCCCCAAGC	TGGTCTTAAA	CTTGGGCTCA	AAAGATCTTC	CCGCCTCAGC
119401 119461	ALCUAGAGTA	GCTGGGATTA	CAGGGGAGCC	CCACTGCGCC	GGCTTGGACT	TTTTTTAATTT
119401	AAACTIGICC	TCTTCTACAT	CTGGTTTTCA	TAACCTGAAG	GCTGTGTTTA	TTTTCCATAA
119521	AACAAGGCAT	IGATICCAAA	GGTATTATAA	TTCCCCAATT	CCGTATAACC	TTCAGCTCTT
119641	TUCCHUMANA TUCCHUMANA	AAAAAAAAA	AAAAAAGAGG	GAATACTGCT	CACCTCCTCT	CCGGAAATGT
119701	ACCCTTTACG	ACTICTUTE	AAACCTTTCA	CAAGAATTGG	ATTUCTTTGT	AATGCTTTAA
119761	TACTCAGAAA	AGTGTTATTG	CACACCETORE	AGCATCTCAA	ACATAGTAGG	ATTACACTAT
119821	TACTCAGAAA	CHILITICIAL	TOTAL CONTRACTOR	ACCOUNT	ATGUTCTTTG	AATCUTAAAC
	TTGCAGCGTT	CIGCHGCTTT	IGITITUTAA	AGCCTAGGTG	TACTCTGCCA	GTCACAAAAT

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119881	GGCGTTTCTC	CAGCACTGCC	GCCAGGTACC	ACCAGCTGGG	AGTTGTTCCT	CTTGCGGAGC
119941	AGGAGGTGGA	CTTGGCCCAA	GAGAAACTGG	ATAGTGGTTC	GCAAGGAACA	TAATTTAGCA
120001		CTAATGCAAT				
120061	CTTTTTAAAT	TCACAAGAGA	CAGGCCACAT	TCTATCTTTT	GATTGGTTTA	GGCTATTTTC
120121	TTGAACAGCC	ATTTAGAAAG	CAGATCTATC	ATCCTTCATT	TGCATGGAGC	GTTCCCATTT
120181	TATTTGAAAC	CAGTTTAACC	CAATAGAAAA	AAGGGAGGCA	GAACCCATTA	TTTAAAGTGG
120241						TTTTCAGATA
120301	CCTCGCTTAT	TACACTAAGA	AAGGTTTATA	TCTTTCACAA	AGGGTTTACT	TACAAAAATC
120361	TTCCAATTTT	GTATACCTGT	GTTTCATAAC	TGACTAGCCG	TCAAACCAAG	ATGTAGAGTT
120421	TCCAACCGTT	ATTTTCCAAA	TTTTTAGAAA	TTACGTGAAA	TATTTGAATG	CATGCCTTCT
120481	CAATAAAATG	GGACGTAGGA	AGCACTGGTG	CAGAAGATGG	GTACAATACT	TATCTGGGAC
120541	CACTCCATTA	TTTGGTTGGC	ACGTTGTTTG	AAGAAAAAGG	GGAAAAGCTC	AGGTTACTTA
120601	GCATGGTTCG	GACTTATTTG	AAAACTACCA	CAGCAGGAGC	GGAAATAAGA	CCGCATTACC
120661	TCACTCTCTG	CTGTGCTGTG	CTAGGGGGTT	ATCCAGAATA	GGATTGTAGA	AGTGGATGTC
120721	GATTTAATAG	TTTTTTTTTC	TCCCATTAGC	TGAGTCTCTG	ATTGGCAATG	TGAGATCGTT
120781	TTAGCTTATT	GATACTTTGA	AATGCACTTA	ACAGCCACAA	ACAAGTTAAA	CCCTTCTTAC
120841	CATAAAATCT	TATCCCCAGG	GTGTGCTTGC	ATTTATCACC	CCTCTTTCCT	TTCACACTAA
120901		CTCCCCAGCA				
120961		GCAGGCTTGT				
121021		TGCGCTGATT				
121081		CATGTTGTAT				
121141	ATTGGACTGT	GGAAATAAGT	TECANETECN	GANACCTTTC	CACACTTCCA	I GCATCCCAC
121201		GTTTGTGAAT				
121261		GACACATCGC				
121321		CACGGCCTAA				
121381		ATTTCTTGCA				
121441	TTCTDATA	ACTCCTCGGA	TECTTOTECE	ACTOCATOTO	GGCAACGGAC	CICIAAAAAI
121501		AAAAGAGCTG				
121561		CTACAATTTC				
121621		GAGGGGAGCA				
121681						
121741	CCACCAATCA	CCTCGGCAGC CGGCGCAGTC	CCACCCCAACA	CGCTAGGGCG	CGAGGGGGGG	GCACTGACGG
121801		TCAGGTTTAT				
121861						
121921	CCCDARCARA	CCCGCCCCCG	CCGCTTCTGC	TGCTCCTGAG	AAACCTTTAG	CTGGCAAGAA
121981	GTCACACCCTC	CCTGCTAAGG	CTGCAGCAGC	CTCCAAGAAA	AAACCCGCTG	GCCCTTCCGT
122041		ATCGTGCAGG				
122101		AAGGCGCTGG				
122161	ACCOMOCCOM	ATTANGAGCC	TGGTAAGCAA	GGGAACGTTG	GTGCAGACAA	AGGGTACCGG
122221	CTCNAACCTC	TCCTTCAAGC GCTACAAAAA	CERROGAR	GGCGTCCTCC	GTGGAAACCA	AGCCCGGCGC
122281						
122341		AAAAAGAGCG AAGAATCCAA				
122401						
122461	CACTGGGAAA	GCTAAGGCTG	TAAAACCCAA	GGCGGCCAAG	GCTAGGGTGA	CGAAGCCAAA
122521	TROTTE COCK	CCCAAGAAAG	CGGCACCCAA	GAAAAAGTAA	ATTCAGTTAG	AAGTTTCTTC
122521	AGRARCECA	ACGGCTCTTT	TAAGAGCCAC	CTACGCATTT	CAGGAAAAGA	GCTGTAGTAC
122581	ACAGATGAAA	TCCCCCCAAGC	AAATGCAACA	CGCCCTCAAT	TATATTAGAA	TCACTTGGAG
	AGTCGATAGA	ACTTTAACAT	AGCCTCATCT	AGTAAGAATT	TACTACTCAA	TCTATCAAAG
122701 122761	ATAGCAAGGT					
122761	GGTAGCCGGA	AGTCCCGCGT	CTCACGACTC	CAAGCTAATT	AGTCATAACC	GTATTGAACC
	AAGGTTGAAG	CCCAGTCCCA	GGCTTGAGGC	TATTATTAT	ACAAGGTTAA	AGTGGGGATA
122881 122941		GGGTCAATAT				
	ATGCTTCTGG					
123001	TAGCGAGCTG	TCTGTCCTTG	GGAAGGACGG	TGACCCTGCT	GGCGTGGCTG	GCGCCCACGT
123061	TGGCGTCCTC	TGAAAGCCCC	GCCAGGTAGG	CCTAGCTCGC	TTGCTTTCTG	CAGCGCCATC

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123121	ATGACAAAGC	TTTGAAACGC	AAAATGCTTT	CTTTGTGCAG	CGCCTTACCA	TGGGTGCACT
123181	TACGGGCTGT	CGACTTGGTT	TAGGCCCTTG	TCAGGACAAA	GGAGCTTAGT	TTGTTGGAGT
123241				GGTTTCTCTG		
123301	CTGATTGGAT	ATTTGAAAAT	TACTGTGCTT	AACTGGATCG	TGTTTCATCA	ATCGTGCAGG
123361	ATTITCAACC	CTGGTGGAGC	CCACACATTC	AAAACTGAAG	ATCCTTTTCT	CAGAACTGCC
123421	CCTTTAAGCT	TTTGCAATTT	TAATTCTGGG	GGTCAGATTT	TAATAATTGG	ACTTTTTTGT
123481	TTACATCTGA	CAAGAGTATA	TGATGAGCCA	AGTTTACTCA	CTTTTACTTA	GTGCAGTTCA
123541	attctaaaag	TTTATTTTTG	CGTGTGTGCA	TATGAGTTAA	TAATCAGTTG	TATTTTTCAA
123601	ACGGTCTTTT	TTCAATTGTT	TTGCTTAGCT	CCTTCCATCG	TCTAAAGTCA	GGGATACAGG
123661	CACATCACAT	CCCTGTTCCC	CCTTCCTCAA	ACTAATATGT	AGCTACCTAG	GTTTATCCTT
123721	TAAAACAAAA	ATTCTCACCT	ATTTTTGTGA	GAAATATACA	TGTTTTTCTT	TGAACTAAGT
123781	ATTTTACATA	CACCTATCTA	TATACATGCA	TACTTGTGGT	TTTGTTTTTT	TAAAAAAAA
123841	AAAAAAAAA	CACGTTATCT	TTTGAGACTG	GGTCTCAGTC	TGTTGCCCAG	ACTGGACTGC
123901	AGTGGCATAA	TCACAGCACA	CTGTAACCTC	CAACTCCTGG	GCTCAGGCTA	TCCTGCAGCC
123961	TCAGCATCCG	GAGTAGCTGG	GATTGCATGC	ACGCACCACC	AAGCCGGGCT	TTTTGTTTTT
124021				TCCAAGCTGG		
124081				GGTCACTGTG		
124141	TGTTTTGTCT	TTTGATTAGG	GTTATTAATT	TAAAAAACAA	AGCCTGGACG	CAGTGGCTCA
124201				GATGGGCAGA		
124261				CATCTTGACA		
124321				CTACTTGGGA		
124381				CAGAGATCGT		
124441				AAATAATAAA		
124501				ACCAAACTTT		
124561				TTATTCATCC		
124621				TGCAATTGAA		
124681				TATTTATTTA		
124741				GCCATGACCT		
124801				TCCCGAGTAG		
124861				GGAGAGATGG		
124921				CTGGCCACCT		
124981				GGACTTTGTC		
125041				ACTGATGATT		
125101				TTTTAAGGCA		
125161				AGCCTTGGAA		
125221				GCTCTCACTG		
125281				ATGGACAGTA		
125341	CTAAGCCATG	GTCAATGCAA	AAGAGTGAGA	AGGAAAGTAG	AATAAGTTAT	CTAAGAATCA
125401	GTGGGTGCTC	TCTTTAAACT	GATTTATCAC	TCCCCCTTCC	AAACTCTCTT	GAAGGTCACT
125461				CTCCAAGGGA		
125521				GGTAAGCATC		
125581	TGTGTTTCTG	TGTTTTTTAT	AGGATCATTC	AATTATTGGT	TGGCTCTTGA	GAGGGAATGC
125641				CTGTATAGAA		
125701	CTAAAATGAG	GCCTGGAGGA	GACATGTTGA	AAGTGACCCA	TAAATCTGCA	GTATCTCATG
125761	TCTCTCAATG	GGGACAAGGA	GTACCATGGG	AAATAGCATT	AGGTCAATGA	CAGTAACAAC
125821	TCCCAGGTGA	GTTGATTTAT	ערדידידים עידידים איניים א	ATAAAGTTGT	TAATATGCTA	CAGIAACAAC
125881	AATTTTGCCA	CAAATAGTCA	TTATTTTAAT	TTCATATTTC	ACTATTGATA	AATGAAGGAA
125941	AAAATGAGTA	GCAGTTAAGC	AGTCCATADA	CCTACATATA	AAGCAAATTG	CACATTTTA
126001	AATTGATTCT	GGATGCTTAA	AATCCTTCTC	ATTGARARA	AATTTCGTAT	TAGAAGATTIAA
126061	CAACATTCTT	TAAACTGAGA	AGCATAACAT	ATAAACAGAA	AACCACAGCA	AAACAAAATT
126121	GCAAAGCTCA	ATAAATGAAC	ACAAAGTGAA	CACCATAATA	ATTGCCACAC	ΔΑςταλαλα
126181	ACAGAAAATC	AGCCAACCCT	CCCAGAGCTG	CCTGATGCTT	GCTTCCAGTC	ביים ביים מיים ביים
126241	TCCATCTGCC	CTAAACATAA	CCCCTATTTT	GATTTCCDAT	CCTCTAATTT	ACTATIONS
126301	TTTTTGAAAC	ATATAAAATG	GAAATAAAAC	AAATGTAATC	CTATCTACCT	CACATATTTC

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126361	ACTCCAGAAC	ATTAGGTTTG	AATAGATTCA	TCTGTGTTGC	TGTGTATAAC	TTTAATTCAT
126421	TTTTATTGTT	ATGTAATATT	CCATGTTATG	AGTGCAACAA	TTTAGGTGTC	TACTGTTGAT
126481						CTGTGTATGT
126541						ATTGTTGGGT
126601						AAATGAAAAT
126661	TAAAACCACA	GTTATAAACA	GCATGGATGA	ACCTCACAAA	CCTAATGTTG	ATGGAATCTA
126721	GCTGGGAATT	CCTGTTCTTC	CATATACTTC	CCAATATTTT	TTTCCAATTA	AAATTGTTAA
126781	TCTTTTGAAG	ATGTTATCCA	TTGTGGCAGA	TGTGCAGTAT	TATCTCATTA	TCCTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT
126841	TTACATCTTT	TGCCCATTTT	TTCTTAATTG	GATTGTATAT	CAGTCGACTT	GGGCTGCCAT
126901						CAGTTCTAAA
126961	GGCCAGGCCA	GAAATCCTAA	ATTGAGGTGC	CAAGAGATTC	AGTTTCTAGT	GAGGGCTCTC
127021	TTATTGACCT	GAAGATAGTT	GCTGTCTTAG	ATTGTTTGGT	CCTCAACACA	ATACCAGAGA
127081	CCAAATAATT	TATABAGAAT	ACAGATTTAT	TTCTTACAA	TOTOCTOCO	ATAAAGCCTA
127141	TEGTCGAGGG	GCCCACCTCT	GCCNACCCC	TTCTTACAAT	TATIOGRAGET	ATAAAGCCTA
127201	ATCTCATATT	CAAACCACAG	CAGTCCCCTT	TTCTTACTGT	TATGGCAGAT	GTGAGATGTC
127261		CCTCATGTCT				
127321		TATGACCTCA				
127381						
127441		AGGCACATTG				
127501		AGATTGTGTT				
127561	CCCTCTCT	TTTTTTTTGA	GGIGGACICI	TGCTGTGTCA	CCCGGGTTGG	AGTGCAGTGG
127621	COCIGICICA	GCTCACTGCA	ACCTCCACCT	CCTGGGTTCA	AGAAATTCTC	CTCCTCCCAA
127621	A COCCOMMO	CTACAGGTGC	ATACCACCGC	GCCCTGCTAA	TTTTTGTATT	TTTGATAGAG
127741	ACGGGGTTTC	ACCATGTTGG	CCAGGCTGGT	CTCAAACTCC	TGACCTCAGG	TGATCCACCT
		CCCAAAATGC				
127801		GTTTTTAAAT				
127861 127921		TAACTTGTCT				
		CCTCCACCTC				
127981		ATGTGCAGGC				
128041		CCAAGTTGGT				
128101		TGGGACTAGA				
128161	ATGCATTAAA	GTCATTAATT	TAGTGTACTC	AAATTAAGCA	CACTGCCCTT	TTATGCACAA
128221		ATCTTATTTA				
128281	TATAATACCT	TCTTGTGAAA	TTAGTTCTCA	AGACTACCCT	CACTTCTAAC	ACCAATTATA
128341		TCTGTGGTTC				
128401		GAAGCTGTTA				
128461		AATGCAATGT				
128521		TCACTCTGAG				
128581	CACATACACA	GCAAAATTCA	TTGTTTGTAA	TAGTTGAAAA	GGGGAAACAA	CTCAAGGAAT
128641		ATCAGCTGAG				
128701		CTTGCCCCCA				
128761	TGTGGCCTTA	TTTGGAAATA	GGATAGTTGC	AGATATAATT	AGTTAAGATG	AGGTTATAGT
128821		GGGCTGGTGA				
128881	GTATTCTTCT	AAGGTGGTCA	CGTGAAGACA	GACACACA	GGCAGAGACT	GCGGTTATGC
128941	AGCTGCAGGT	CAAGGAATGT	CAAAGGTTGC	CAGCAAGTAC	GAGAAGCTAG	GAAGAGTCAA
129001	GGAAGGATTT	TCCTACAGGC	TTCAGTGGAA	GCATAGATCT	AATGATACCT	TCATGTCAGA
129061	TTTCTAGCTT	CCAGAACTAC	AAGAGAATAT	ATTTGTTGTT	TTAAGCCACC	CTAGCTTCTA
129121	GCTCTTTGTT	ACAGCAGCCC	TAGGAAACTA	ATATAGGCAC	AATCCAGGCA	AGTTCCAAAT
129181	ATGAGCTTCC	AGTTGTCCTC	TCCCAGTAAT	ATGAACAGTA	TTACTTTCCC	AGCATTAATG
129241	TGTGACAATA	CACATGACGT	ACAGAGCAGT	CCCCACTTAT	GCACAAAACA	TATGTTCCAG
129301	GACCTCCAGT	GGATGTCTGA	AACCATGGAT	AGTACTGAAC	TCTATATAGC	TGTTTTTTCC
129361	TATACAGACA	CAGCTATGAT	AAGGCTTAAT	TTATAAATTA	GGCACAGTAA	GAGATTAATA
129421	ACAATAAATT	AGAATAATTG	TTAAGAATAT	ACTGTATAAA	AGTTAGGTGA	ATGTTTATTT
129481	CTGAAATTTA	CCGTTTATTA	TTTTTGGACT	GCAGTAGACC	ACAGGAACTA	AAACCATGTA
129541	GAAACCGTAT	ACAAGAGAAC	TGTATTTCAC	CCGAGCCTCA	GTGTGCAGTT	TTAATGGCCT

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129601		ACTGCTCACA				
129661		CAAAGTTCCT		_		
129721	CCAGGTAAAC	AAAGACACAC	TTATCAGTGA	GAACATTTCA	AGGGTCTAAA	ATTCATCTCC
129781		GGGCAAAGGC				
129841		TTTTCATGTT				
129901		ATAGTGTGAA				
129961		AAGGCTATAC				
130021		GTGCCACTTT				
130081		TATTTTTGGA				
130141	TTTAATAAAG	CAACAGATTT	GTTTATTTTG	GGCCCTTGGA	TTTGTGTATT	AAATTTGAAC
130201	CCTGTTTGTC	AATTTCTATA	ATAAAGCTTA	TTGGGAATCT	GATTAGGATT	ACAATGGTTT
130261	TGTAGATCAG	TTTGGGGACA	ATTAATACCT	TTAAAATATT	GACCGCTTCA	ACTGTAAATA
130321	TACTCCTCCA	TTATTTAGTT	TTCCTGTTTA	ATTTATCTGA	GTAATACATT	ATAGTTTTCT
130381	TCGTAGAAGT	CAGATACGTA	GAAAATTCAA	AGCCCAAGTG	CAATAGCTCA	TGTCTGTAAT
130441	ACCAGCACTT	TGGGAGGCCG	ATGTGGGTGG	ATCACCTGAG	GTCAGGAGTT	TGAGACCAGA
130501	CTGGCCAACA	TGGTGAAACC	TCATCTCTAG	TAAAAATACA	AAAATTAGCT	GGGTGTGGTG
130561		GTAATCCCAG				
130621		TTGCAGTGAG				
130681		CTCAAAAAAA				
130741	AATAACTAAA	AAATGAACAG	TTATTAAAAT	ATCAGGATAT	AAAAGCAAAA	AAATCAATAA
130801		TACAAAATGG				
130861		CTCCCTGAAA				
130921	GCTTCATGCC	CAGATATGGG	TACAGTTCCT	TGTTTAAAAA	AATTTTGCTC	CATCAATCAA
130981		CTTCCTCAGA		•		
131041		TAAGGAAGTT				
131101		GCACAACTGC				-
131161		CAAGTCCGTG				
131221		GACCTCCCTG				
131281		ATTCCTTTTC				
131341		GCATTTATTG				
131401		ACTATCCACC				
131461		TGCATGTATT				•
131521		CTTTTGCTAC				
131581		CTGAGGCAGA		_		- <del>-</del>
131641		TAGAGCAAGA				
131701		AGGTAGAATC				
131761		TAACTGGAGC				
131821		TTGTCTCAAA				
131881		ATGGTGCAGG				
131941		AATCATGAGT				
132001		TGGTAACTCT				
132061		ACACTTTTGT				
132121		CTCTTGTGGC				
132181		AAAACAAAGG				
132241		AAGGTAATTC				
132301		TTAGTCAACA				
132361		AGCAAAGGTC				
132421		GGAAGAGGAG				
132481		GAGGAGGAGA			-	
132541		AGAAGGAGGA				
132601		ATATAGGCAT				
132661		AACACCATCA				
132721		GACAAAAGTT				
132781		AACTAAAAAG				
	AUGUMON1	Shunurian	GCACICIGGA	TITUGHMAIN	GGWWGICWIT	VG T CWCCT T C

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132841		AGCCAGAGGA				
132901		GGAGGTGTAA				
132961	CAGCTGCATG	GAGATTTGGG	AGAGGGAAAG	CTTTTTTTT	TTTTTTTTAA	TTGGAAAAGA
133021		TGTAAATAGA				
133081	AAAAACAAGT	GCACAGTTAT	CTAAGGGAAA	CAATGGGATC	AAGCTGCAAG	TATATAAACT
133141	TGTCTTGATA	GAAGAATCCT	TGATCTGGTT	TATTCAGTGT	TTGGTCCAAA	CCCACATCCC
133201	TGTTCTGCCT	GTCTCTGACT	TGCTCTGTGC	CCCAGAAGCC	CAGCTTCTAC	AGATAGCATT
133261	AGCTGGGCAG	CCCTGCCCTC	TTGCAACAGC	TGGATTTGGC	CAGTGATCAG	CCCAGCAGGA
133321	ATGTAGATGG	CAAAGGAGAG	AGAGGTTAGT	GTACTTATTC	CCTGCATCAC	CCCCCTGCTT
133381	GGTGGGCAGC	TCTTCCTCCA	CAGTCCCAGC	TCTGGCCTAG	CTCTGGTTAC	AGGTTCCCTC
133441	CCATTGCCTC	TTCAGATTTA	AAGGTGTGTC	TGTCAGGGTA	TAACTGGGAG	CTAGAAATTG
133501	CACTGAAATT	GAACAAAGAA	TTTTATGGGA	ATGGTTGTTA	ACTAGTTATA	AGAGGACTGA
133561	AAATGGAAAA	GTGGAACAAA	CGTATCAGAG	ATAGTAATGA	CAGAAAGCAA	CTACCACCTC
133621	CAGGTTTAGG	AGAACAAGGA	AAAGATTCTT	TGAAGAGATC	CCCAGAACTG	GGACCTCTGA
133681	GGAGTGTATG	CTGGACCACT	GATGATGATA	TGTCTGTAGA	TAGAGGCATG	ATGAGGCTGA
133741	TTTTAGGAGC	ATGGAAGATC	TCCAAACTGA	AGCCAACTGC	TGTTACTGGA	TTCAACTGCC
133801	ACTGCCAGGT	TGAAGAACCC	ATTCTGTGAG	GATGTCAACA	AACAAAGTGG	GAAATCTTTT
133861	CACATCCTTC	CAGCCCTCTA	GTCTTCCTCC	AGTGCTTTCT	ATTGGTAGGG	TTTGGGGAGG
133921	TGGCTAGCAA	AGCGGTATTG	GAAAAGATAG	AAGAGACTAA	ATCTTCATAA	CCAGCACAGG
133981	GTGACACTGG	ATCACTACTG	TTGCTGATCT	TGGGCTGCCT	CATATCCCCT	GTTCTTCCCA
134041	TTAGCCCTGT	CACAACTTTG	TAGATATCCC	TTCATTATAT	GCCCTTCATA	TATTCTTTTG
134101	GTTTAACTTT	TTCTGTTGGA	ATCCTAATAT	GGCACTCCTC	CATTTTTCAG	GACCAAAAGA
134161		TTATCTTTTA				
134221		ATCTATACAT				
134281		AAAAATAAAA				
134341		CACGATGGCT				
134401		TTATAGGGGT				
134461		AATAGCATGC				
134521		TCTACACTTA				
134581		GGAAACAGAA				
134641		AAAATCTGTG				
134701		GTCATGCTCA				
134761		CCCATAATCG				
134821		AGACACAGAA				
134881		GAAGTGGCAA				
134941		GTTAAAGCTA				
135001		ATTTTCCACA				
135061		GAGCAAAAAT				
135121		AATCTAAGTG				
135181		TTCGCAACAC				
135241		AGTTACGAGT				
135301		GCTTGGGCGC		-		
135361		TAGTCTGATA				
135421		TCCTGAAATG				
135481		TGCCTTGTGG				
135541		CATCTTCCAG				
135601		AAAGAAAGAA				
135661		GAGGTGGGAG				
135721		AAGACCCTGT				
135781		AAAGAAAGAG				
135841		GTTCAGGATC				
135901		CATCATCTCT				
135961		CATTGTCTCT				
136021		GCACCTAGAG				

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136081				AGACTTGCTA		
136141				CAATGTCAGG		
136201	AGAGTATGGA	AACCCAATGG	GAAAAATGCT	CACCAGGAAC	ATGAAGAGAA	GGAATTACGT
136261	GTAAGGATTT	CTCAATGTGT	TCCCAAATTT	GCCCAGCAGA	GGGAGGCCTC	GGGTTGATGG
136321	CAGGCTGACC	ACACAATTAA	AGAAGGCTGA	ACCTGGGGGC	TTTTAACAAC	CATCGTGGGC
136381				ATCCATTCAA		
136441				ACTTTTCTAC		
136501				ATATTCAATT		
136561				CATAATAATC		
136621				GTCACACTCT		
136681				ACCTCCTGGA		
136741				TGCACCACCA		
136801	TTTTAGTAAA	GACGGGGTTT	CACCATGTTG	GTCAGGCTGG	TCTCCAACTC	CTGACCTCAT
136861	GATCTGCCCA	CCTTGGCCTC	CCAAAGTGCT	GGGATAATCA	CTTTTTATGC	TGCATAATTC
136921				CTCATTTGTT		
136981				TTAATATGAT		
137041				TAATGTAATT		
137101				TTATAAAGCT		
137161				TTACAAGTAG		
137221				TAAAAAACAA		
137281				TAACAGAACC		
137341				TAAAAGTCCC		
137401				TTTTCTGAGA		
137461				TCACTGCAAC		
137521				TGGGACCACA		
137581				GGTTTCACCG		
137641				GCCTCCCAAA		
137701				CCAAAGAAAA		
137761				TGACACATTT		
137821				TTAATTCATA		
137881				CAGCAGAAAG		
137941				GAAACCCTTC		
138001				CATGTTAGTG		
138061				TGGCAAAAGT		
138121				AGCCAATGTG		
138181				AAACTCCTCA		
138241				GGTGGGCAGA		
138301				CATCTCTACA		
138361				TACTCAAGAG		
138421				AGATCACGCC		
138481				TCTCCTCGAA		
138541				AAAATATTAA		
138601	CCACACCACA	CCGTATCCCA	CAGGCACTGC	GGATGTGCAT	ATGCAAGAGG	GGTTGATCCT
138661	AAGAATTTAG	AGTCACAGAG	GAGGAGGCAC	CAAGCAGACT	GTGGAGAAAG	TCATGACCAG
138721				TATCTGGCCT		
138781				TTAGGTTTCT		
138841	GATCGTTGAC	CCAGTTAGCA	TTCAAGCAAC	TTCCACCCTG	CACTTTTATT	CTTTCCCCTT
138901				TAATAATAAA		
138961				TCTAACTCCT		
139021				CAGGTAGTTT		
139081	AATAACTATA	AAATATGGTA	AGTTTATAAA	AGTTACATTG	AGTATACTTT	ATAAGAACTG
139141	CTTATTGAGT	TTGCCTAATA	ACCACACAGC	ACAATAATAA	TATGTATATA	TTTTTAAATA
139201				AAGGTATATC		
139261				TGGCATAGCT		

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139321		CAAAAAAATA				
139381		AGGGTTGGGC				
139441		GAAAAGAAAG				
139501		GTGTTCAGCA				
139561	ATTGATCATT	ATTGGGTGTT	TCTCGGAGAG	GGGGATGTGG	CAGGGTCAAA	GGATAATAGT
139621	GGAGAGAAGG	TCAGCAGGTA	AACACGTGAA	CAAAGGTCTC	TGCATCATAA	ACAAGGTAAA
139681	GAATTAAGTG	CTGTGCTTTA	GATATGCATA	CACATAAACA	TCTCAATGAC	TTGAAGAGCA
139741	GTATTGCTGC	CAGCATGTCC	CACCTCCAGC	CCTAAGGCAG	TTTTCCCCTA	TCTCAGTAGA
139801	TGGAATATAC	AATCGGGTTT	TACACTGAGA	CATTCCATTG	CCCAGGGACG	AGCAGGAGAC
139861	AGATGCCTTC	CTCTTGTCTC	AACTGCAAAG	AGGCGTTCCT	TCCTCTTTTA	CTAATCCTCC
139921	TCAGCACAGA	CCCTTTACGG	GTGTCGGGCT	GGGGGACGGT	CAGGTCTTTC	CCTTCCCACG
139981	AGGCCACATT	TCAGACTATC	ACATGGGGAG	AAACCTTGGA	CAATACCTGG	CTTTCCTAGG
140041	CAGAGGTCCC	TGTGGCCTTC	CTCAGTGTTT	TGTGTCCCTG	AGTACTTGAG	ATTAGGGAGT
140101	GGAGATGACT	CTTAACGAGC	ATGCTGCCTT	CAAGCATTTC	TTTAACAAAG	CACATCTTGC
140161	ACAGCCCTTA	ATCCATTTAA	CCCTGAGTTG	ACACAGCATA	TGTCTCAGGG	AGCACAGGGT
140221	TGGGGCTAGG	GTTAGATTAA	CAGCATCTCA	AGGCAGAAGA	ATTTTTCTTA	GTACAGAACA
140281	AAATGGAGTC	TCCTATGTCT	ACTTCTTTCT	ACACAGACAC	AGTAACAATG	TGATCTCTCT
140341	CTCTTTTCCC	CACAGGAGGT	GATGGCCGGA	AGAACATGGC	AGAGGGCAAA	ACAAAACAGC
140401	ATTGGGAACA	AGCTCTGTTT	AAAAGGAGAC	TTGTGAACAG	CAAAGAGTAG	AAAGGGTTCT
140461	CTTACAACTG	AAGCCCATGG	AAGACAAATG	TGTACTGCGT	GAGTTTTAAG	GCAATAGGAG
140521	TAGTGGGACC	TAGGGCACAC	CAGAGAGCAT	ATTAACTCTC	AAACTTTTAA	AAACATTATA
140581	TCTGCTGGAC	ACAGTGGCTC	ACACCTTAAT	CCTACAACTT	TGGGAGGCCG	AGGCGGGCGG
140641	GTGTAGCTTG	AGCCCAGGAG	TTCGAGACCA	ACCTGGGCAA	CATGGCAAAA	TCCCGTCCCT
140701		CAAACAAAAA				
140761		AGGCTGAGGT				
140821		ATGCCACTGC				
140881		ACACACCATA				
140941		ACTACTAAAT				
141001		TCTTAGGCTA				
141061		TAAGTACTGT				
141121		TCAACAAAAT				
141181		TGTAGTTGGC				
141241		CTGGCATGCC				
141301		TGCAGCACCG				
141361		AGGCTGCTCT				
141421		TTTACCCATG				
141481		GGAGAATAGA				
141541		TCTGCAAATC				
141601		ATCCTGTGGC				
141661		AATCCAAGGA				
141721		GACCAGGTTAT				
141781						
141841		AGCCATAGAC				
141901		CTGTAGCAAA				
		GGCCTGTCTC				
141961		TTGGGATATT				
142021		AATGATGTAC				
142081		TAACTAAGGT				
142141		TCATTTGTAG				
142201		CAGAAAGTTA				
142261		AGTTTATCTC				
142321		GAAAGAGATG				
142381		TCTAGTGTTC				
142441		AGCTAGAAAA				
142501	CTTGAAATAA	TGGATATTCT	AATTAATTAC	CCTGATCTGA	TCACTATACA	CAGTATGTAT

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142561	AAAAATAACA	CTATGGGCTG	GGCGCAGTGG	CTCACACCTG	TAATCCCAGC	ACTTTGGGAG
142621	GCCAAGGTAA	GCAGATCACT	TGAGGTCAGG	AGTTAGAGAC	CAGTCTGGCC	AACATAGTGA
142681	AACTCCATCC	CTACTAAAAA	TACAAAAATC	AGCCAGGCGT	GGTGGCATGT	GCCTGTAATC
142741	CCAGCTACTC	AGGAGGCTGA	GGCAAGAGAA	TTGCTTGAAC	CCAGGAGGCG	GAGGTTGCAG
142801	TGAGCCGAAA	TCGCGCCACT	GCACTCCAGC	CTGGGTAACA	GAGCAAGGCT	CTGTTTCAAA
142861	AATAAATAAA	TACATAAATA	AATATTTTTT	AAAAAAAGAA	CATCACTATG	CACCCCATAT
142921	ATACATATAA	TTATTATGTC	AATTTGAAAC	ATAATTTTGA	AAAATGAAAA	AATGAAACAC
142981	AAATATGAAT	CAATCCTCTC	CAAGTTGATA	TACTTAAAAG	GAAAAAAGTC	CGAGGGCTTA
143041	AACTATTCAA	TCAAAATTTT	ATTAAAATGC	TATAGTAATC	TGGAAAGTAT	TTCAGAATGA
143101	ATTGGTATAA	GGTTAGACAC	AAAGATCAGT	GAAACAAAAT	AGAGAACCCA	GAAATAGATT
143161		TGGACAACTG				
143221	ATCGTCTTTT	CAGTAAATGT	TTCTTGAACA	AGTAGACATC	CGGTGTGGGG	GAGAGGAGCA
143281		CTCAAACTTT				
143341		ATTATAAAAC				
143401	TAGCAAAGAT	TTCTTTAAAA	CAAAACAACA	GGTTTATAGT	TTATAAAACA	TAAATAACAA
143461		TTCATCAAAA				
143521		AGGCATGAGA				
143581		TGCACTCCAG				
143641		AAATAGAAAA				
143701		GACAAAGGAC				
143761		AAAACAAAGA				
143821		TATGCACATG				
143881		AATGAGATAC				
143941		TGAGCAAGGG				·
144001		TATAACTGAA				
144061		CGAATGGTTT				
144121		CACGTCCTTT				
144181		GGGTATCTTA				
144241		GAGTCGGCCT				
144301		TCCTCGCGAG				
144361		TCCAGCTGCA				
144421		GTGACTGACG				
144481		GTGCTGAAGT				
144541		ATTGACACCT				
144601		GATACCGCCT				
144661		GCCGGGCATC				_
144721		ATTCACATAC				
144781		TAATGACCAA				
144841		TCATGCCTTT				
144901		AGAGATTAAT				
144961		ATATAAATTA				
145121		ACACCTTACA				
145181		GACAAATGGT				
145241		TGTACCACAG				
145301		CAGTGTCCAA				
145361		TGTCCTCAGG				
145421		AAAGAGAAGG				
145481		ACCCGATGCC				
145541		GAAAAACAGA				
145601		CACTTTTCTG				
145661		GAAATAAATA				
145721		GAGTTACAAG				_
145781		ATTTTTATTT				
145841		CTAATTGTTA				
	<b></b>					

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145901	GTAGAAGCGA	GGCATGGTGG	CTCAAGCCTG	TAATCCCAAC	ACTTTGGGAG	GCTAAGGTGG
145961	GAGGATTGCT	TGAGCCCAGT	AGTTCAAGAC	CAGCCTGGGC	AACATGGAGA	AACCCTGTCT
146021		AAATGAGCCA				
146081	GCTGAGGTGG	GAGGATGACT	TGAGCCTAGG	CAGTCAAGGC	TGCAGTGAGC	CCTGATCTTG
146141		CCAGTCTGGG				
146201		AAAATGAAAG				
146261		AGAACAATAA				
146321		GCAATTGCAA				
146381		CCCTTTGGAA				
146441		CAACATAGTG				
146501		AGTAGACAAT				
146561		TTCCAACAGT				
146621		AAAAAATATT				
146681		TTAAATTTTT				
146741		ATTTCAGTTA				
146801		AAAGTGAAAG				
146861		ACTTATGCCT				
146921		GAGTTCGAGA				
146981		AAAAAAAAAG				
147041		AAAACTGTAA				
147101		TTTGGGTGCA				
147161		CCACACATTG				
147221		ATATACAAAG				
147281		TGACCTGAAC				
147341		GCTTGACATC				
147401		AGTAGAATGA				
147461		TAGTAACTTG				
147521		GGTGGTTTCT				
147581						
147641		GTATTTATCC				
147701		TGGATAACTG				
147761						
147821		CACTTTGGGA			<del>-</del>	
147881		CAACATGGGA				
147941		CGGGAGGCTG				
148001		GCCACTGCAC				
148061		AAAAAAAAA				
148121		AGCCATGAAA				
148181		ACTGCATACT CCGGGAAGAA				
148241						
148301		CAACAGCAGA GAGTAGCAGC				
148361		AGGGGGATTA				
148421		AAGAAGTCGA				
148481		GCGTGTCTTA				
148541		CCGGAGTAAA				
148601		ACTCTAGAAA				
148661						
148721		ACAGGAAGTG				
148781		TATAATGGTG GAATAAACCC				
148841		ATTGTAATAA				
148901		ATTAGTTACA				
148961		ATTCAATGTG				
149921						
149021		CCTAGCAGAT				
74300T	AGAMAGACC	AGAAAGCTAG	CICICITIT	GCCATGTGAA	GACATAGCAG	GAAGG TAGCC

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149141	ATCTCCA ACC	TAGGAAAGGG	COMMORCANA	CARTCARCTC	ACACCTCACA	N C N C T C N C N C
149201		GTTGTTTAAG				
149261		TTGGATTAGA				
149321		TTAAGCTTTT				
149381		TTAAATAGTA				
149441	TTCTCTGTTT	TTTAATATGA	GAACTCCTGA	TTTTCAGCCA	ACTACCCTGG	AAAAAAAGCT
149501	AATCTTTCTG	ACTTCTTAAG	TGTGGCCATG	TACTAAATTC	TGGCTAATGC	AAGGCAAGCC
149561	AAAGGTTTTA	TGATAGGTTT	TAGGACACTA	GAGTAAAAGA	GAGCTGTTGC	ACACATGCTC
149621	TTCACCCTAC	TTTTGTGTCC	TTTTTTCCAT	CCTACAACTT	GGGTTGTGAG	TATGATGGCT
149681	GGAACTTTAG	TGGCTCTCTT	GGATCCCAGG	GGTAATTGAG	GGGTGGCTGG	AAGGAATCTG
149741	TGATTTTCTG	GAGTTTCCAT	ACACAAACAA	GACCTGGATT	TTCTGGGCTT	CCCAGACTTC
149801	CACATCTAGA	CTTGCTTTAA	ATGGGAGAGA	AATAAACTTG	TTTCAGCCAC	TGTCATTTTG
149861	GGCTATTTTA	TAGAACTTAA	TCTAATCTTC	AAGGGTACAT	GAATTGCTTT	TCCTTAAAAA
149921		CATAAAATCA				
149981		TTTTTTTTT				
150041		CATGACCATG				
150101		CCTGAGTAGC				
150161		GATGGGAGCC				
150221		GCCTCCCAAA				
150221		CTGAATTTCA				
150341						
•		AGGCCATTGG				
150401		TCTCAAAATT				
150461		AATAGCTAAT				
150521		TTAATCTCCA				
150581		TCATTTTTTA				
150641		TCTAAGTGCT				
150701		TAGGGAGGCC				
150761		CCAAGAGGTA				•
150821		GGTGGGGTTC				
150881		ACTACGGGTA				
150941		TTTTTTTCTA				•
151001	TAACAAAAGA	CAGATTGACA	AGACAGAAAC	AAAGCATGTG	CATTGTACAA	ACACAGGGGA
151061	GTACTGAGAT	GAATACTCAA	AAGAGGATTT	AGAACTTGGG	CTTATATAGC	ATTTTAAGAA
151121	AAGAATACAT	TTTTTAAGTG	ACAAGGAAGA	CGAAAAGGAC	TTTGAGTTTC	TAGTGCAGTA
151181	AATTGTGGGA	AGGCAACTTT	TTCTTTCCCT	TTTTTTTTT	TTTTTTTTTA	AAAAAAAGAC
151241	TTCTCTGGTG	CTATGTCCAG	GCTGATAAGA	GTCTAAAGTC	TCTGGTGACT	AACTTTTGTT
151301	CTTCCCCGAG	TAAGAAGACA	CCTTCACAAT	TTCATATCCT	GCTTTTAGGC	AAACAGGGAG
151361	AGGGCAGAGG	TGTTTGTTTG	TTTTTAATCT	ATTTTTTTC	TCAATTGTCT	TCAACTCAAA
151421	ATACTTCTTA	TGCCAAAGAT	GGCATATTCT	GCTACCCTTC	ACTTACTACT	TACAACCCAG
151481	CCTCTATCAT	CATAATTAGA	ACTTCTGACC	CTGGGGAACA	TGGGCAATAG	TTTGAACTCT
151541	TTTATATCTC	CCTTAGGCAG	AGATGGAGGC	CCAGCCATGC	CTCTGACATC	TAGACACAAC
151601	TGTTGCTTCA	TTTCTCCTAT	TCTCAGAGGT	GATGTTGTAG	GACTTCAACA	AATATCAGTA
151661	AACATTAATT	TTTTTTTTCC	TTGAGGCACA	GCATGATCTT	GGCTTACTGC	AGCTGCTGCA
151721	GGCTCAAGCA	ATTCTCCTGC	CTTGGCCTCA	CGAGTAGCTG	GGTTACAGGC	CCCTACCACC
151781	ATGCCCGGCT	AATTTTTGTA	TTTTTAGTAG	AGACAGGGTT	TCACCATGTT	GGCCAGGCTG
151841	GTGTTGAACT	CCTGACCTCA	AGTGATCCAC	CTGCCTCAGC	CTCACATAGT	TCTGGGATTA
151901		CCACCATGCC				
151961		TGACTTTTTA				
152021		ACTCGGGTAT				
152081		GAATCATAAT				
152141		ATATTTCAAG				
152201		ATCCCCCATC				
152261		AAGCCAGTCT				
152321						AACATGTCAG
<b></b>			CCICCAGGAC	- CCCCCI IAG		

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152381		AGTCAATGTT				
152441		GCAGCTTTCT				
152501		GACCTTCTGA				
152561	ATCAAAGCCC	TGACCCATTA	AAGACGGATG	GAGACAGCAA	CATACGATCA	TCACTATTAT
152621		CCCAGTCCAG				
152681		ATCAATTATA				
152741	ATGACAGCGA	AGAAAACAGA	CCAAACGTCT	GCCCTTATGG	GATTTATATT	ATTTTCTCTG
152801	TGCTGGTTAA	ACCAAGGAGC	TTCTGCTCTT	TTCCTTAGTC	ACCTGGGGGA	GGCAGAAACA
152861	AAGGAGAATA	TTGATAAACC	TGGAAATAGG	GCCGGAGAGT	ATCAGAGAAG	GAAGCCTTCG
152921	GGAAAGTAAA	GATGTGGCAG	CCAGTATTCC	CGTTATAAAA	GGATACAACT	CCGGCCTCAT
152981	AGTCCAGAAA	AATTCCCACA	AGCAGGGGCT	GCTCATGCAG	ATGAAGGGAA	GTTGGGGGAG
153041	AAGTAAGTGC	TACATAGCCT	TTCTTTTTGC	ACAGCCTGAG	GGTCCAGAAT	CCAGACTGAG
153101	GCTCTTGCTT	CATGCCAGTG	CCCCTCTGCA	CATTTTCCAT	ACAAACTCCT	AAATCCCATC
153161		GCCAACATCC				
153221		GGGGAAGGCA				
153281		CACTTGTCTC				
153341		CACACTAACT				
153401		AGGTTATTTT				
153461		ACTGGCACTA				
153521		TCAGCCTCCG				
153581		AATTTTCTGT				
153641		GAAGTGTTCC				
153701		GCCCAGCCAC				
153761		TATAAACTGG				
153821		AAAGTGGAAA				
153881		TTGCTTAATG				
153941		GGGGTGATCA				
154001		TGAGAAAAGA				
154061		GCTGCAAAGG				
154121		AGGTGGTGAG				
154181		GTGGTGCATT				
154241		TCTGCTGCTT				
154301		TGTGCACTTT				
154361		CACAAGGGGT				
154421						TGGACCCTGG
154481		TCTTCCAAGA				
154541		ACATGTAGTA				
154601		GGTACAGAAC				
154661		GACTTCCAGG				
154721		AACCACCAAC				
154781		TGGAAGACCC				
154841		TCTTTCCCCT				
154901		AATATACATG				
154961		AAAGTGAGAT				
155021		ACAAGAGAGT				
155081	GGTCTGTGAC	AATTAAGGAA	TTCAAGTATT	CAGAAACACT	TCCTGTGCTG	CATCCTCTCT
155141		CCAAATAATC				
155201		ATTAACCAGG				
155261		ATAGGGCACA				
155321		GAGGGTCTGC				
155381		GCCACTTGAG				
155441		TGTTCCTTGT				
155501		TTCAGTTAGC				
155561		AATGACCCAG				
					~~ <del>~~~~</del>	

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155621	AATACCACTT	AAGAGAAAA	ATATCAATTG	GATTTTTAAA	ATTCCACCTA	TCTATTGGTG
155681	TGACACATCA	ACAAAAACAT	ATAGAAAGAT	TGGAAGCTAA	AAGATAGATA	ATATAGTCAT
155741	ATACTGTTAT	AGTATTATAT	CAAAAGATAT	TAAGTCAGAG	CATTATTAAG	AATGGAAGAA
155801						AGGCAGGCGG
155861						CTGGCTCTAC
155921						CTACTTGGGA
155981						CTGAGATTTC
156041						AAAAAAAAGA
156101	AAGAATGAAA	GGAGTCACCT	AAAAAAGATA	ACACAATTTT	AAACATAAAT	GTACTACATT
156161	ATTAGTGAAT	TCATGTTTAG	AATTGTGTTA	ATATACAAAG	CAAAAATTGT	AGAATTATAG
156221	GAGAAATGGA	CAAATCTACA	ATCATCATGG	GATGTTTTAA	CATTCTTCTT	TCCATAATTG
156281						GAACAATCTA
156341	AGAAGCGCAA	TCTCATAGTC	AATACATAAA	GCTCAGCAAT	TGTTTAATAA	TAGTAAGCAG
156401	AGAATATGCA	GTTTTCTCAG	GTATAGATGG	AACATGCACT	AACTGAGTAA	ATACTAGGCA
156461	GAAAACAGTC	TGAACAAGTT	TCAATAAATC	TGTATTACAC	AGATCATTT	CTCTAGCCA
156521	AATATAAGAT	TATAAACCAA	TAATAAAAAG	ATGACTAAAA	AGATTCTAAA	TATTAGGAAA
156581				TATAGAATGG		
156641	TATAAAAATA	TACAATGAAG	CTABAGCAGA	ATTTTAAGGA	ANATOTOTAL	COMMUNICATION
156701	CTTATCTTAG	מממידממממ	DAGCTCAACA	TTAATGAGCC	AMAITIGIAG	GCITTAAATG
156761	AAAAAGAACA	TAGAAAGCCA	AAGCIGAACA	TTTAAAAAGA	AMGCAICIMA	TITAAATTT
156821				CAATAAAAAA		
156881						ATTTATCAAA
156941				CAGAAACTTT		
157001				AAGTACTGAT		
157061				AACAAATAAA		
157121				CCGACAGATA		
157181				GAAATTTAGA		
157241						TAGACATAGA
157301				AACAAACCAA		
157361				AACATATACA		
157421				GTATTTTACA		
157481				ATGACATATA		
157541	GTGTACAATA	ATCAGGCCAG	CCACACTCAC	TCATGCCTGT	TARIACATIA	CATCAAAAAT
157601	CTGAGGCGGG	TCDATCACTT	GACACAGIGAC	GTTTGAGACC	ARICCCAGCA	CGTTGGGAGG
157661				ATCCAGGCAT		
157721				TCACTTAAGC		
157781				CTGGGTGGCA		
157841				GCCTGTTCCT		
157901				GTGGTGGAGC		
157961	ACTCGGCTTG	GGCAACAGAG	TCACACCCTC	TCTCGAAAAA	CATGATCGCA	CCACTGTACC
158021	ATCAATATAT	ביית מיית מיית מיית	TROCARTORA	TGCTTCACTT	WCWWWINIWI.	ACACACAGTA
158081	ATCTTATTAG	ATATATACTA	TACCARICAG	ATAGATAGAT	ACATACACAM	TATAGATTAC
158141				TATATATATC		
158201	TCTCAAAAA	משתותותות	CACCCACATA	TGGTGGCCCA	TATAGCATAT	AGAGATGCTG
158261	GGGGAGGCTG	ADATCACACC	ATTCCCAMIG	ATCCTCTCAT	TGCCTGTAGT	CCCAGCTACT
158321	TACTGCACCA	CTCACCCTCC	CACACACACA	CACACTCAT	TGGTTGAGCC	ATAATCGCAC
158381	ΔΤΔΤΔΤΑΝΑΤ	CICAGCCIGG	ACACACAGAGG	GAGACCTGAG TAATATATGT	G TGGAAGGAT	ATAGATATAG
158441	ATTATCANCA	CACTGGGAGA	CARTACTATA	TATATATGT	GIGTATGTGT	ATATATATAT
158501	GAAGACACTC	GTGGGATGGT	THE TAKEN CON	ATTGGACCAA	CACTGCCACC	TATATATTAT
158561	CATGCAATGT	TCTTCTTCAC	TONITACEA	GAGCACTGGT	CAMACOMIA	AIGGAGCCAA
158621	GTCTCCAATC	AGACATACTT	ADDOLIGGER AND TATE	ATGAACTTGC	CATAGITACG	CACACAMOMO
158681	GTGTGTATAT	VCCCATACTI	CACCAACCTA	GCAATTGTCT	CHIMIACGIG	GAGAGITUIG
158741	TCAGAGCADA	GATGACAGCT	VACCAWCCIN	GCARTTGTCT	TCATCATCAT	IATAATGCTA
158801	CTCCCCCACC		COMCOMCOMO.	CTTCATCT	CONCORD	CITCUTTCCC
<del></del>				CATCATCTCT	CITCITTTT	IIIIIGAGAT

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158861	GGAGTCTTAC	TCTGTCGCTC	AAGCTGGAGT	GCAGTGGCAC	AATCTCAGCT	CACTGCAACC
158921	TCTGCCTTCT	GGGTTCAAGC	AATTCTGCCT	AAGCCTCCAG	AGTAGCTAGG	ACTGCAAGTG
158981	CACACCACCA	CACCTGGCTA	ATTTTTGTAT	TTTTAGTAGA	GATAGGGTTT	CACAATGCTG
159041	GCCAGGCTGG	TCTCAAACTC	CTGCCCTCAA	GTGATCCTCC	TGCCTCGGCC	TCCCAATGTG
159101	CTGGGATTAC	AGGCGTAAGC	CACTGTACCC	GGCCTCCTCC	TTTAATAGAC	AGGGTCTAGC
159161	TCTGTTGCCC	AGGCTGGGTA	CAGTGGCGTG	ATCATAGCTT	ACTGCAGCCT	CGAACTCCTG
159221	GGCTCAGGAG	ATCCTCCTGC	CCTAGTCTCC	CCAGTAGCTG	GAACTACAGG	CATAGCACAC
159281	GGGGCTAATA	AAATTAATTA	GGTGATAAAA	TTCACTGCCC	ACTGATGACT	AAGCTCTTTG
159341	GACATAAAAG	ACACAGACCT	TGAAGGAAAA	TGTGTCTACT	TAATTTCAA	ACCCTATTTA
159401	TCAAAAAACA	GGATGAAAAT	GCAAAATGCC	ATCCACATEC	CAGAAGATAT	CAGCTATAAT
159461	AAGTTCCCAT	AAATCAATAA	GGAAAAGAAC	CCADTANAN	TTATTATA	ACAGTAAATC
159521	ATGGGTAAAT	CACAGAGGCC	TGAAGGGCTA	ATGGACATAC	AAAAAAACC	TCAATCTCAC
159581	TAGTGAAATC	AGAAAAGCAC	AAATTAAGTA	CACAATTAGG	TACCAMMMA	1 CAATCTCAC
159641	ACTGTCAAAA	TCATAAATTA	TATAAGTAAA	GACTCAGGGA	GTTTTGGACC	AATCTGTAAG
159701	CTTATATTGC	TTGTGGGGTA	GAATTGGAAC	DATTTONACA	TOTOTACTAC	AGTGAGAGCT
159761	TATGATATGC	ATCCCTCACA	CCAGCATGTC	DOTCOBROAT	ATCTCCCTCC	CIGGIAAAAT
159821	TACGGGACAC	AAGGAAGCAT	GGATAAGAAT	GTTCACAGGI	GTATTCTCTCTC	AGGGAACATT
159881	AACAACAAAA	AAACCCAACT	ACACACAACT	TCDATCCCCA	CTCCACAAGC	CAACAGCAAC
159941	AATAAACTTC	AGGCCGGAGA	TEGTEGTTCA	TCARIGCCCA	CCCALAGG	CAATGGATTA
160001	AGGCGAGAGG	ACTGCTTGAG	CCCAGGACTT	CARCACCACC	CCCAACACTT	TAGAAGGCCG
160061	GTGTTTCTAC	AAAAAATTTT	TARARARTA	CCCACACCAGC	CIGAACAAAA	TAAAGAGATA
160121	CAGCTACTGG	GGAAGCTGAC	GTGGGAGGAT	TCCTTA A CCC	GCAGTGCTTG	CCTGTGGTCC
160181	GAGCCATGAT	GGGGCCATTG	CACTCCACCC	TCCCTCAGCC	CAGGAATTTA	AGGCTGCAGG
160241	AGATAAGTAA	ATAACAACTT	TCCATCCAGCC	CCCACATTTCC	AGTGAGACCC	TGTCTAAAAG
160301	TCTAGACTCT	AGACTCTTTC	TATCACTACC	GCCACATIGC	AAAATGGTGA	GAGAGTGGTT
160361	TAACCTCTCT	GTGTCATATT	TOTOTOTOTA	TICIAGITAT	AMGGGGGGA	AACACTCACC
160421	TGTGATATAA	AACAAGAACC	ANGARANGTA	AAAGCAAAA	ATGCCCCATA	TAGAGAGGAC
160481	GTGCTCAGTA	TATGTGAGTC	AAGAAAAGIA	AAGCTTTTCT	AATCTGTCAC	AGACTAAAGA
160541	AGTCAAGTAA	TATGGTACCA	TATIALICCIG	GIGCIGGIAG	GAGTGTATGT	TACAACTITG
160601	GGTATGTTCC	CARAGGIACCA	CARACCACCA	TTAACAACAA	CCTCGGCAAT	CCCAGTTTGG
160661	TTGTAGCAAC	CAAAAGAAAT	GAAAGCACCA	GGATATAAGG	ATGCATGGAC	TAGAAAGTTA
160721	ATGGTTACAT	ATTGTAATAA	CIAAGIICIA	AAAACAGCCT	GAAGCTCCAT	CAGTAGGGAT
160781	ATAGAAGAGA	ATATTTATTA	PACTED COME	GAATATTAGA	CATAAAAAGT	AACGAGTAAC
160841	CCTACCTAGA	CAGTGTATAT GAAGTCAGAT	TCCACACCCC	TGTACAAACT	TAGGGAAAGA	TATAGATCAC
160901	CCTACCTAGA	CTTTCACTCA	TTTAAAAGGGG	TGGGAAAAAC	CTTGAACTTT	CTCCTTATAT
160961	מתחתמממדם	GTTTGACTGA	TIAAAATGIA	TTTGTTGCAT	CTGCTTGAAG	GCAATGTAAA
161021	AGCTGGGGAG	CATACATTTA	AAAATAAAAA	TAAAATTTAT	TCCTATCACT	TTTGTAATAA
161081	TCACCTCACC	AGTGACTAAC	ACTIGIAATC	CTAGCACTTT	GGGAGGCAGA	GACAGGCAGA
161141	LOUCETONGG	TCAGGGGTTT	GAGACCAGCC	TGGCCAACAT	TGTGAAACCC	CATCTCTACT
161201	GGCTGAGGCC	AAATCAGCCA	GGCATAGTGG	TGCGTACCTG	TAATCCCACG	CTACCCGGGA
161261	AGCTAGCCTG	CTGGAACCCA	GGAGGCAGAG	GCTGCAGTGA	GCTGAGATTG	CGGCACTGCA
161321	TAATAAACAG	GGTAACAGCG	AGACTCCATC	TCAAAAAAA	ATTTGAAAAA	AGAAAAATTT
161381	TAGTETICAG	TGTTTAAGAG	GGGAGAAATA	TITAGTTAAA	AGATAAGCCC	ATTTAAGAAA
161441	CAGCCTCCCC	TGACCCGGAA	GGCGGAGCTT	GCAGTGAGCC	GAGATCGCAC	CACTGCACTC
161501	AAGAAATAGT	GACAGAGCGA	GACTETETET	CAAAAAAAA	AAAAAAGAAA	GAAAGAAAGA
161561	AAGAAATAGT	TCACTIGAA	CCATATTATG	ATTCCTTCTG	TAAAAGATGA	GAGTAGGCAA
161621	ATTGACTCAG	CARAMICCCA	GCAAAACTTA	CACAAAGTCT	TGTTCTTCCT	TCCTGTCATC
161681	TGTATAGGAT	GAAATACAGA	GIGCITITGG	GITTTGTTGT	TGTTTGTTGT	TGTGTATTTG
161741	AGGGGAACAC	AGGTCTATAA	TICCTITICT	GAAATCCCTG	GAACAAAATG	GGCTTTGCCA
161801	TTCAAATTAG	ARCCCACAC	ATAAAGGCAA	AAAAATGCAT .	ATACTCTAAA	GTTCAACCCC
161861	ATCATGGCCT	AAGGCAGAGC	CCTGTAATCA	AATTCATCAA	TATATCTGCA	GCAAAACATT
161921	TATTCAAATT	MAGIGGGATA .	AATAAAGACT	TTTAAATAGT	CTCATCTCAG	TGCCGTTCAG
161981	GGTTGGCCAC	CCTANARGAC	AGACTCAAGG	GIGGCCITCT .	ATGATTCCTG	CCTCTTGGTG
162041	TTCACACCCT	CGIAMAATTC	CTTGTCTTTG	AGTGTGAGCA	GGGCTTATGA	ATTGCTTCTG
	ACCAATAGGA	INIGGCAAAG	ATGATGGGAT	ATAATITCTA '	TGATTACGTT	TCATTATGTA

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162101						CTCTCTGAAG
162161	AAATAACTGG	CCATGTTAGA	AGCCCATGTG	CAAAGAGCTG	AGGGGTGGCC	TGTAGAAGCT
162221						AACCATCAGG
162281	AAAGAAATTC	TGCCTGCTAT	CTCAGTGAGC	TTGGAAGTGG	ATTCTTCCTT	AGCCTAGCCT
162341	CCAGATAAGA	ACACAGCCTG	ACCAACACCT	TAACTGCAGC	CTTATCAGAC	CCTAAGCAGC
162401	AGGCCCAACT	AAGCTGTGCC	CAGATTCCTG	AACCACAAAA	ATTGAGATAA	CATATCAGTG
162361	TTGTATTAAG	GTTCTAAATT	ATGGTAATTT	GTTTGTACTA	ATAGATAACT	AATATAACCA
162421		TCAGGTTAGG				
162481	ATTTTCAGGG	GTTTTTTTGA	TTTTGTACTT	ACGGATACAA	ATTTGTGAAA	GTATAGTCAG
162541	CACTGATTTA	AAAAATCAAG	GGAGCAGGAA	ACTCAGTAAA	TGGTTCTAAC	ATTTTGGAAT
162601	CTGTAAATTG	GTTGTAACAT	TTGTCATCTG	TGTTATCTAA	GTCAAGTTCC	TAAAATATGT
162661	GAATGATAGG	TTATCATACT	CACCTACTTT	TCTTGCATTG	CTCTAAGAGT	TGGCTGAGCT
162721	ATTGATAATA	AACACTATGA	TCAGATCTAA	TACCATGATG	TGCTATTATG	ATCATGTGTC
162781	AGTCACAGGG	CTAAGCACTT	TGTACATGTT	GATGCATTTA	ATTTTGATGA	TAACTCAATG
162841	AAGTAGGAGC	TGTTAATATT	TTCATTTTTC	AGAGGGGGAA	ACCAAGTCAC	TTGGAGTAAC
162901	ATGGCTAATA	AGTGAAAGAA	TAAGAATTTG	AAAGGTTTGC	ACAGATAACC	AGAATGCAAT
162961	GCTCATCACA	TTCACTGAGC	AGTGAATCAT	ACTAACTAGA	GAAAGTATGA	ADGCTCTACT
163021	GAAATTAACT	AAACAACCTC	TCTGGCTGTG	AGCCTGCCAA	GGGACAGGTG	CTANACTCIACI
163081		AGGCCCCTTC				
163141	ATGACTCCTT	AACATTTTCT	TCACATCGAA	GTAAAGCTTG	CDAACATTCC	ACATACCITO
163201	AAGTTCCAAG	GAGACAGCCT	CTGATGTTTC	CAGCTTCACA	GCCCAACTCC	TAGAATAAGC
163261	AGAGGCGAGA	GATTTCTTCA	GAGGTGCATT	CCATTCATTT	CTATATACCC	ACACCCCTCC
163321	CCTCCTGCAT	TCAAACAGGA	CTTACCTCCT	Chargeges	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	ACACCCICC
163381		GGTGAGCATG				
163441		TTTACCCGAA				
163501		AAACAAAGTA				
163561		CCAGCTTCTG				
163621		CTTGCTGAAG				
163681		GGTGATCGGA				
163741		AGTGGTGAAT				
163801		GTATCACACA				
163861		CCTGTTAGTT				
163921	GATAAAGCTG	TAGATTGTAC	CATARCAGAR	CTCCACCAAC	COMMONACA	TCTTCCTACA
163981	GAAAAGGTTA	TCACTCATTT	TOCOTOGOG	A COUNCE OF CARGO	GGTTCTACAA	GCAAATCAGG
164041	TATTTCACCA	GAGAGTCAAC	AACCACCCCC	ACTICATOAC	CCATCAGTCA	CCTAGTGGAG
164101						
164161		ATCCCGTGGT				
164221	ATTOTALA	TAAAGGTACC	CTATAAATTA	GTGAAGGCCA	GCCTGATGGC	ACTGATGTAC
164281	TATAACATAC	AACATTACTT	TATCTTCCCA	TGCTTCCTTA	CCATTCTCCT	TTAATAGCAC
164341		CTTTTTTCCC				
164401		CATTTTTCCT				
164461		GCCTCATAGT				
164521		TAAGACTTCT				
164581	CTCCAACTAA	TGTCTCTGAA	TCTGTACCTT	CTCCTGGAGT	CAAGAAAGTA	TGGTCAAAAG
164641	JCJCCCTT CT	ACCAAATGTC	CATCTATGGA	TGAATGGATA	AACAAGAATG	AAAGTCTGAC
164701	CCARAMAN	ACATGACAAG	CCTTGAAGAC	ATTCAAGCAA	AATAAGCCAG	AAACAAAAGG
164761		TAAGACTTTG				
164821		TAGTGGTTAC				
164821		AGTTTCAGTT				
		TGGTTGCACA				
164941 165001		AGTGGTTAAA				
165061		AGCCACTGGG				
		TGCTGCAGAG				
165121	AAGGREGGE	CTTATAGTCA	CCCAATTAAT	TITCTTGTTC	TTCAGCCAAG	ACACAGGAGA
165181	AAGCTGGGTT	AGGAGTGCTA	GATAATTTAA	TTGTGAAACT	AGGGCCAAGT	TCAAACACTT

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165241	TATCAGTTAC	AAGGATAAAA	AGAGGTTTTT	ACTTATGATT	TAAGAAGTTA	GATTTCTGAG
165301	TTGGAGCGAT	TTTCTTGAAG	TAAAAGCTTA	TAATGAACAT	CACCCAGACT	GGATTTTAAG
165361	ACAACCAGGC	TGGTAAGAGG	GTCCATAATT	CTTGGCAGGG	GGAGCTTTGA	GTGTGACAGG
165421	CATTTATTAT	GGTTAACTGA	GAAATACTGT	TCTACTACCC	TAGGGTCATC	TTAAGCATTC
165481	CTATGTGTAA	GACTGACAGA	AATCAAGTGA	AACTCTCATC	TGAGGAGATG	TAAAGTTGCA
165541	ATTTCCATTA	GTGCTGTCTA	AATTAATGCA	GTGGGAGTGT	GTATTCAGGG	CAATTTGAAT
165601	CTATGTTCTT	GGATTGCAGT	CTTCAAACTT	GGCCCAAATA	AACTCTCTAC	TTATCTTAAA
165661	AAAATAAAA	TTAAAAAATA	AAAATAAATT	CATACAGTGT	TTTGATGACT	ATGATATAGA
165721	AGAAGGGTCT	TTGACTTAGG	ATGAGGTGGA	ATTTTTGTGT	AGGAGACAGG	TGCAGCTTTA
165781	ACTCTTGTAT	AGACGGGTTT	TCATATATGT	TAGTTACAAT	CAAGGTCTTC	CCCATTGCCC
165841	AAGATCCTAG	AAATGGGGGA	AGTAAGAGTG	TACTCAGGAG	CTCAAGAGCA	ACATCCACAA
165901	ACAAAGATCA	GGGTAGAGGT	TAGAGAGGAC	TCCTGAAAGA	GAGAAAATTG	GTAATCAGCT
165961	TGTGGGATTT	TACTGCAAGC	TAGTGAATTA	TATAAATATA	AAGATTGGTG	CAAAAGTAAT
166021	TGTGGTTTTT	GCCTTTACTT	TAATGGCAAA	GACCGCAATT	ACTITITICCAC	AAACCTAAAT
166081	ATTTCCATAA	AAGAATGTGG	CTCTGATAAT	GTGGAGGTTA	GTCAGCCACG	CABATAATCT
166141	GAAAGTTTGT	AGTTGCAAGT	GTGTAGGTTG	TTGCATTACT	TGTGATGTAC	TTATAAATCA
166201	AGTATAGGCC	GGGTGCAGTG	GCTCACGCCT	GTAATCCCAG	Chemmeres	CCCECACOEC
166261	GGTGAATCAC	GAGGTCAGGA	GATCAAGACC	ATCCTGGCCA	ACATGGTGAA	ACCCCCTCTC
166321	TACTAAAATA	CAAAAAATTA	GCCAGGCATG	GTAGCACATG	CCTCTAATCC	CACCUCATOR
166381	AGAGGCTGAG	GCAGGGGAAT	TGCTTGAACC	CGGGAGGTGG	ACATTECACT	CAGCIACICA
166441	CGCACCACTA	CACTCCAGCA	AGACTCCATC	TCAAAAAATA	CTANTANTO	GAGCIGAGAT
166501	AAATAAATAA	AGTATATTTC	TTTCATCAGC	TTCATCACCT	TCACTACTAT	CAAMMOOCAAM
166561	CTGGAGTGAT	CCTGTTTTCT	AAGTGTTCAC	AAACCCTCCCT	TOAGIAGIAI	GAATTICAAT
166621		CTCCACTGTG				
166681	GTTTATTTC	AGGTATTTAA	ACTOTO	CAGGGTAATG	AGTTGAGGCC	TGCAAACCAG
166741	ATACTAATTC	TGCTTCTTCT	GAGGAC	CCACTCGATG	CTTTTTCTAG	GTAAATAGTC
166801	ATGGAAAGAT	TCCTCCTAAA	GAC IGAAGIA	TCAGGAATCC	CAGCCAACTA	CAGTTTAAAG
166861		TGGTGCTAAA				
166921		TCTTCCTTGG TCCTCAATCA				
166981						
167041		CAGATTCTTA				
167101	DAGGARART	CTGTCTTCAA	DACCOTTCAG	TTTTGTCACA	GCTTTCTGGA	GCTITTCCTG
167161	AATCCCAATC	TTGATAAGTG	MAGGCCTATTC	AATTTGACTC	TTCATTAGGG	ACCTAGGGGG
167221	TATECCARIC	TTCTAAGATA	TATTTGAATA	ATAGTGAATA	TTTATAGAGT	CCTCATTGTT
167281	CCTCAATACT	GAGCATGCTA	AAGGCTATAT	GTGCAGGAAC	ATACTGATCC	CCTTGGCAAC
167341	ACCCCTARA	TGGTAGGATT	TTAAACTTCA	TTTCTGTGCT	GTAGAAAATG	AGACTAAGAA
167401		TAACTTGCCC				
167461	TTCCATCIGC	TGACCCAGAG	CCTGAGCTAT	GTCCACCACT	AGAGTCCTGC	CAGGAAAAAG
167521	ACCARARCO	AACAAGGTAA	TCATCATCTA	AAAGATTTTG	TAAAACAACA	TGCTGAACCA
167581		ATACCAGTGT				
167641		CAGCTGGTTA				
167701	DOLLARDAME I	TATCATGAAT	CCAATTTAAA	ATGATTTAGT	ATTCATGTCA	AGCTTTTAGC
167761	COULTMAN	AAACAGTTTC	TCATATTTCT	ATTGAAAGTG	ATTTGAAGCT	GACCCAAATT
167821	GCTAATTGTA	GTCAATGCTG	AAAGAATTGT	CTCCTGTCCT	CTGTAAACCC	AACAAGTATA
167881	CTCATTCATT	CTCGAGTGTT	CTCAGGAAAA	GGTTCTATGT	AACTGTTTTA	GCAAAAGATG
	ACATTGTCCT	TACTATATGC	CAAGTGCTAT	TCTATGCATT	CTATATTTTA	ATGTCCTCAA
167941	AGCTTATAAC					
168001	TTTACAGATG	GAGAAACCAA	GGTGTGAAGA	CATTAAGTAA	CGTGCCCAAA	ATTGCCCATC
168061	TAGTAAGTGA	CAAAACTCAA	TITCAACATA	AGCTGGTTCC	TTTTCTTACT	ACTTGGTGGA
168121	AAAGTAATTC	AAATGGGAAT	ATGATCATCG	CAGTTATTAG	CTGCTCCATG	GAGTTTAAGG
168181	AAGAGCTGCC	ATGAGCTGAG	TGGTGGTCAT	GATTGACATG	TCCTTAGAAG	GACTTAGAGC
168241	CTTCATACAA					
168301	AACATTTTCC					
168361	TTCCTCCATG	CTGCCAACAG	CAAAGTCCCA	CCTTCCTTAA	TATGCTTTCT	GGCAAGAAAT
168421	CTGGATGGTA	CACAAAACCT	CTCCCTCTGC	TTCACCTTCC	ACAACCAAGC	ATTTCCAAAT

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160401	CTTTTC 3 CTTCT	mommoomes s				
168481 168541	CITIGACICT	TUTTUUTGAA	TCGTGCTTAA	AATCTGCCCT	CTCCTCCCTT	TCTTATACGG
	ATAGTTTGAA	TTTTACTCCT	TGATATTCCT	TTTATCATAG	ACATGCCACA	GTAGCTGGGC
168601	ACAGIGGIIC	AIGCCTCTAA	TCCCAGCATT	TIGGGAGGCT	GAGATGGGAG	GGAGACCAGG
168661						TAAAAAAATT
168721						GGTGGGAGGA
168781	TTGCTTGAGC	CCCAGAAGGT	TGAGGCTGCA	GTGAGCCGAG	ATTGCACCAT	TGTACTCCAA
168841	CCTGGGATAC	AGAGCAAGAC	CCTACCTCAG	AAAAAAAAA	AAAAAAAAA	AAAGTAGAGG
168901	TACCAGAGTG	ATATTTTCAA	TGTCACTGAC	CCTTCATTCC	CCAAATGAAA	ATCCCCCAAT
168961	AGGTGTTCAA	TTTTTACGTG	TCCTTCAGGA	GTTACTTCTA	AGATGAACCA	CTCTCTACCC
169021	TAAATGTCCC	TCCCCACCAC	CAAAACCAGG	GACCTCCAGG	CAGACATTTT	TGATGGTTTG
169081	TTTTCTTTAC	TAGACTGTAG	ATACCTAAAA	GGTGATGGGT	CTTTCTTCCC	TGTTTTCAGG
169141	CCCTACTGCA	TGGCTTTACA	TATTGTGGTT	TTTCAAATGA	TATTCATGGT	GTGAAACAAG
169201	AAAAAATGCG	GGTGTTTGGT	TTGAGAACAA	CCTGTTCTAA	AGCAAAAAGA	AATTCATCAT
169261	AACACAAATG	GATAGAGATA	AGAGTCCAAC	CATCCCATTG	AAGGTCAGGA	TGGACAGTCT
169321	AGATAATTGA	GCAAGAAATC	ATCATAAACT	ATTTTTCAGA	AGAATGACAT	GATGAAAGCT
169381	GTATTTCCAA	GTCATAATGT	TAGGTTTCAA	GTTAAATCAT	CTCAGCTCCT	GGGGAGCAGG
169441	ATAAGACTTG	GTACTTACCA	AAGCTCCCGG	GCCCACACAC	TCACCTTGTA	GCCCTGGCAT
169501	ACGTCTTCAA	CAAGAGCTGT	GGTGTGCCCT	TTGTGCTGTG	GTGCCCGCTC	ACAGCGCCAG
169561	CAGATGAGCT	GCCCCTCATC	TTCGCAGAAC	AGGTGGAACT	GCTCTCCGTG	TTCCTCACAT
169621		GATCCGTCTC				
169681	CGGAGGCTAT	CCATATGAAA	TGGAGCCCGA	CACTGGGGAC	AGCAGAATGT	CTCCTGCCTC
169741	AGTTGCTTTT	GGCTTGGGTT	TTTAAAGAAG	TCTGTTATAC	ACAAGTGGCA	GTAGCTGTGT
169801	CCACAGTTGA	TGCTTACTGG	GTTCGTCATC	AGGCTCAGGC	AGATGGAGCA	GGTGGCTTCC
169861	TCCATCATCT	TCTTGGTGCT	GGTGGTTGAG	GCCATAGCTT	TTATTGAAAA	GCTCCAATAT
169921	TGGCTCTAGA	GATGGAGATG	AAGCAGCCAG	AATTTTCCAC	CGTGATGAAA	ATACACCTCA
169981	CCTGCACCTC	TATGTGATGA	GCTGGCTGCA	ACTGACTTCC	ATAGGTCTTG	AAGGTTTTCC
170041		TATTATCTCA				
170101		TGTTTGGGCC				
170161		CAAGCAGTTA				
170221	TTTTAAAGTT	GTTTGCCAAG	AATTTACATT	AAAATAGCAT	AAGCTTTTGA	CTGGCTATAC
170281		GTATTACAAA				
170341		CTTTTAAACA				
170401		TTTGCATACC				
170461	CTTTTCTCAG	TCTTCTAACT	TTTTTTTTTT	TTTTTAATGA	GACGGAGTCT	CACTCTGTCA
170521	CCCAGGCTGG	AGTGCAGTGA	CGCTATCTCG	GCTCACTGCA	CCTCCGCCTC	CCGGGTTCAA
170581	GCGATTCTCC	TGCCTCAGCC	TCCCGAGTAG	TAGCTGGGTC	TACAGGTGTG	CACCACTACG
170641	CCCAGCTAAT	TTTTGTATTT	TTAGTAGAGA	TGGGGTTTCA	CCATGTTGGT	TGGCTAGGAT
170701	GGTCTCGATC	TCTCGACCTT	GTGATCCACC	CGCCTCAGCC	TCCCAAAGTG	CCAGGATTAC
170761	AGGCATGAGC	CACCGTGCCC	AGCCTCTTTT	TCTTTTCTTA	TAAGACAAGT	TCTCGCTCTC
170821	TTGCCCAGGC	TGTAGTGGAG	GGCAGTGGCA	TGACCACAGC	TCACTGCAGC	CTCGACCTCC
170881	TGGGTTTAAG	CAATCCTCCT	GCCTCACCCT	GGCAGAGTGG	CTGGGACTAC	AGGTATGTGC
170941	CACCATGTCC	AGCTAAAGTC	TTCTCTCCAG	AAAGAAGAAA	TGCATTGGAA	TTTAGAGGAT
171001	ACACAAACAT	CTAGCTGTAT	AGCTAATACA	GTAGCCACTA	TCATGAGTAG	GAATTTAAAT
171061	TTAACTTAAT	AAAAATTAAA	ATGAAAAAAT	TCAGTTTTTC	TGTTCCAGTT	GCCACATTTT
171121	GATTGCTTAA	TAGTTGCATG	TGACTAGTGG	CTACATAACA	GCCTCAATAT	ACAACATTCT
171181	GTTATCACAG	AAAGTTACCT	TGGACCAAGT	GCTGGGAGAA	GCAATGCAGG	CTTCCTCACA
171241	AAAGCTGTAA	AAGAGAGAAC	TCAGGGAGTG	TGAAACTCTT	TCCTATTCTA	GTTAACTTCA
171301	AGAATAATTG	TTACCAGGCC	AGCACGGTGG	CTCACGCCTG	TAATCCTAGC	ACTTTGGGAA
171361	GCCGAGGCGG	GCAGATCACC	TGAGGTCAGG	AGTTTGAGAC	CAGCCTGACC	AACATGGCAA
171421	AACCTCATCT	CTACTAAAAA	TACAAAAAGT	TAGCTAGATG	TGGTGGTGCA	CACCTGTAAT
171481	CCCAGCTGCT	CAGGAGGCTG	AGGAAGGAGA	ATGACTTGAG	CTCCGGAGGG	GGAGGTTGCA
171541	GTGAGCCCAG	ATTACACCAC	TGCACTCCAG	CCTGGGTGAA	AGAGCGAGAA	TCTGTCTTAA
171601	AAAAAAAAA	AAAAGAATAA	TTGGTACCAG	AATTACTCTT	TGTAATTAGT	AGTAACACTT
171661	ATGCAATTGG	GTGATCTGTG	ACAGATTCCA	TTGAAGGAGT	ATGGGGAGCT	TCACCCCAAT

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171721	ATATGACTCC	CTGGTATAAT	GAGTATTTT	AATTAAAGG	CCTTAGAGAT	CAGCAGATGC
171781	TGGAAGAGAC	: TITTCCCCT	L TCTACATAAA	GACCAGTCAC	ACTAGACAAC	ב ממממממת
171841	GTTTTTCCTT	CCAACCCCTA	TTATCTCATI	TTGTACTGA	GAAAAGAGG	CTAAGAATGT
171901	AACCAGACCI	: AATCAGACAC	: TTTCACAAAA	TAATGTCTGT	CTCTCAGGCT	יייייייייייייייייייייייייייייייייייייי
171961	CCAAAGAGAA	CCATTTACA	GTTAAACTCT	GTTCCTCCA1	TCATTCATCC	TCCCSSSTAT
172021	TCATTTATTC	TCCCTAGTA	TCATTTACTG	CCCCTCAAAC	AATTACCTAT	ATTCTCCTGA
172081	TATCACCCTI	CCCCTCTGAA	ATAAATATGT	ATACATGTAT	ADACGTTATA	CATACATATT
172141	TATACAGTAT	ACATACATAT	TTATACATAC	ATACATATGO	' ATACATATTT	ATATTTATGT
172201	ATTTATACAT	: AAGTATTTAI	' AAATAAGGCT	ATATAAGTAT	CTACCCCCAT	TOCONCROS
172261	GGTAATCACT	CTGTGATTCT	' AGCCCATGTA	CTTGTTAATA	AATTTGTATG	CCTTTTTCTCC
172321	AATTAGCCTG	CCTTTTGTGA	GTCGATTTTT	CAGTGAACTT	CAGAAGGCAA	ACCCCA ACTC
172381	TICCCTTGGC	TCCTACACCA	. TCATGACAAT	AAAATTTGAC	TCCACCTCCA	CCCCCCCCAT
172441	CCCCCACACA	GAACAACAAC	CAACACTGGT	TAATAAGGTC	CCT.LCT.T.T.	The state of the s
172501	TTTGTTGTTG	TTGTTGTTGT	'TGTTGTTTTT	GCTTTCAGGA	GCAGAGGTAT	AATTACCCAAA
172561	AGAAAGAGAA	AGGAGAATAG	TGAATACCTC	TTCTGCAGAG	AGGGGTGCCT	ABGTGGGACT
172621	TCCCTGGCTA	ATAACGTCTT	GCTAGAGACC	CAACCAGGAG	GATAATGGAA	GCAATCAAGG
172681	CAACCAGAAC	AACCAGAAGA	ACCAGTTTAT	CCTTTTTGTG	CCCTCTCCCT	AAACTGAGGG
172741	AATAAGAATT	GGAAAGAAGG	CTGCAGAGCA	GAGGGTTTGC	TCCTGAGGAG	CAGTTATTTC
172801	TATGGGATCA	GAGCTCCTGC	AGAACTGGGG	AGTTTACTT	TACTATCTCT	TCTCCAGGAC
172861	AGGACCTATC	TCAAGAGACA	TGTTCAGAGT	GATTGCAACA	TABAGAGTTT	GCAGACCCAA
172921	GGAGGTAGGG	AAGGCAGAAA	GAAGATGGGG	GAGGCCAGGG	ATAGGCAACA	GAGGAGTGAC
172981	CAGGAGCGAA	AAAGCCTGCC	TCTTCTGAGA	ACCTAGCTGG	GCTCTCCCTG	TACCCCCGAT
173041	CCCTCCCCC	CGCCCGCCCC	CACACCCCTA	CTCCTGGGAG	CTCCTCTAGG	ACAGGGGCAG
173101	AGTCAGGAGG	AAGTTTGAAG	AGTGCCTAGA	ATAAAAAAACA	GTAATTTAAC	TACAATTACC
173161	GGGTAGGCTG	TTTTCCTCTC	ACAATTTGAT	CAGTCTCTTG	AAGCCACACA	CAATHITICATIC
173221	TGAAGACGTG	TATTCCTTGG	CAGGCTATTT	CCTCCAGTGA	TACACCAGGC	CCCTCTCTGC
173281	TGGGGTCACT	GCTCTTCTGG	GGAGATGGGG	CTCCCCTCCT	TCCAAGGCTC	CAGGGTTCCT
173341	GTCCTGGGCC	CCACTCATCT	AAGTTCTGAA	TCTTCTGAGA	TTTGGTGTAA	AGTCTGGTGA
173401	AAGAAAGAGC	AGGAAAGAGG	TGAGAGCTGT	AAAACAAAGA	AAGTCCTGAC	CATTTTCAGA
173461	GTTGGAGGGG	CCCTGCTGTC	ACGAAATATA	TTCCCCACCC	CACTTGCCAT	CAGTACACAC
173521	TCACATATCC	ACTGAGAAAA	CCTTAGCCTG	GACCTTTTCC	GTAACCTTCA	CTGCTCAGAC
173581	ACTTACATAT	TCGCTGCTAG	TCCCCTCTGT	TGCTGCCACT	TCCTGGGTCA	GGAAGTTAAC
173641	TCAGACCGGA	TTAAACTGAG	AAGTGAAACT	ACTGTGGGAG	GCGGGGCTCA	ТАВСВТОТАС
173701	GAGAAAACTA	GTGACGTTGT	TCATATCATT	TGCACTCCGC	CTCTCCGGTA	AAGGAGGGG
173761	AAACGTAGGA	AGAAAATATC	CTTCTTTTAC	AGCAATAAAA	AGAAGGAACC	ם מתדמ מדדמ מ
173821	CCTGTAAACT	ATCATGTGAC	CCCAACACAG	AGTATCTAAA	AACAGGAAGC	CTGCAGAGGT
173881	TCAGTTCACA	GACTCTGATT	TGAGATCTTT	CTACTTTTGC	CACCAACTCC	CTTGGGAGTC
173941	CTTAAGCCTT	CCTAGCTGAT	GTTACTTCTT	TTGCTATTTA	TGGGTTGCTT	GTGGTTCTDT
174001	AACTGCTCTG	AAGGGTGTGG	TGGAAAAAGG	GGTGGTAACA	GCAGTAGGAC	TCATTGGCAT
174061	CACAAAATTC	ATCTGAGTCA	GCTTTCTATT	CTTCTCTGTC	CCGTTCTGTG	<b>Վուսաբարարի</b>
174121	CICCIIGCIG	TCCTTCTGCA	GGACTCAGAT	CTTCTTCAAT	AGCGAGGGTC	ACCCACCATA
174181	GAMAATGGGA	GTCACTAGTG	GCCCAGCAGT	GAGTGCCCCC	AGCTTAGAGC	TOTOTOCOAT
174241	CCCIGGGACC	ATCACTCTGC	TTTGTGCTTT	GTGGAGAAAA	GGCTGTGGGG	TCCAGGGTCA
174301	AGICCITAAT	GACTTAGCTC	CAGCTTCTCC	ACTTCAAAAT	GAAAGGAAAA	GTACTATCAC
174361	CACCCGTTAG	AATTATTATT	TCATGGGGAA	AAAAGATGGA	TTACTATCTC	ACAATAAGAG
174421	CITGTCACAT	TTATAAGTCT	CAGGTGTAAG	AGGCATTTAT	GATAACAACA	<b>ፕልልጥልአልፕሮ</b> ሮ
174481	IGGCTTAAGT	AGATGCAGTG	GTCCAAGGGA	ACCAGTAAGG	GGAGCTCAGG	ACACAGGTGG
174541	GAGGAGAAAT	TAAACTTGAA	TTCTGGGAGC	CACTGGCCTG	TCTGGGCCCC	<b>TGGCCTGCCT</b>
174601	GCIGACCCTG	ATAGCCAATG	GAACATGGAG	TTTGGCCCAG	CTGCAATCCC	TCTGGTCCAA
174661	CIACICAAAA	TAAAGGCAAG	ATTGGGAAAC	ACGTTCCTTT	CTTCCTATAC	CAAGCAGAAG
174721	ACTUTTUAGE	ACTGCACCCT	CCTGGGTGCT	CACAGAGCCT	TCTGTTGTTT	ጥርርሮልርርሞልር
174781	GATTCATCAT	GCCCTGGCAT	GATGGTTGCA	GACCCCATGC	ATAGCATGGG	<b>ACATTCTACT</b>
174841	CCIGAGGCAA	CCAGCACACA	GAGAGAGGAG .	AAAGAATGAG	CCCCTGAATC	СТТССТСССА
174901	CGATGAGTCC	TTGCAGATAT	CTACAACTTT	CATTGTTGTG	GATGTGACTC	TGTACCCAGG

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174961	CATGGCTCAT	TCCAGATCTG	TCCTATTGTC	AGAGGTGTTC	AAACCAGAAT	GACTCCATTT
175021						AGGAAGTTAG
175081						TAAAGATCTT
175141						AAGATGGCAA
175201						GTGTTAGCAA
175261						ATGTCAGGAA
175321						CCCACATCTT
175381	TCCCAGAAAA	CATATGAATA	ATCCACTCCT	TGTTTAGTAC	ATAATCAAGA	AATAACTGTA
175441	AGTATCTGTA	TTAGTCCATT	TTCACACTGC	TGATCCAGAC	ATACCTGAGA	CTGAGTAATT
175501	TATACCAGGA	AAAAATGTTT	CATGCTCTTA	CAGTCCCACG	TGTCTGGGGA	GACCTCACAA
175561						AGGCAAAGAG
175621						ATTGACTATC
175681		CAGTATAAAT				
175741		CTCCCACAAT				
175801		AGTATCCTTA				
175861		ACTTTCTTGC				
175921		AAGAACCTTC				
175981	TCAAAGGATC	AGGAAAAGGA	AGCTAGTGAA	TGCTAAAAAG	GAAACAAACT	ACCATTACCA
176041		CAAGACAAAA				
176101		GGAAGGAGGG				
176161		ATGAGACTTC				
176221		CCCAAGGAGG				
176281		ATATAGGAGT				
176341		TGTACTGTGT				
176401		CCTCAACTCC				
176461		GTACCCCTCT				
176521		TGAATATTTT				
176581		TGGATTTTT				
176641		GTGATGTGCA				
176701		GTGTGTGCAC				
176761		GTACTGTGTG				
176821		TGTGGTATGT				
176881		TGGTACTAGA				
176941		CCCACCTGTA				
177001		TACTGGGCTG				
177061		GCCAAGGGAA				
177121		TTTATTAACA				
177181		AAAGCTTATA				
177241		GATGGGATTA				
177301		ACAGCCAGGA				
177361		TGAAAGGAAA				
177421		ACAGTTGAAT				
177481	TCCTACCAGT	GCATGCAGGC	CTTTAGACTG	AGCTACTCCA	TATTGATTAA	TTTCCTGAAC
177541		TTAAGGAAAG				
177601	CAAGTTCCCT	TATCTGCACA	AAACATCCGG	TGTAAGCACT	TGTGGGGCAG	GTCAGAGGTT
177661	CTCTGGGTAC	CATTCCCTTA	CTGTCTGCCT	AAAGCAAGCT	GGCCAACTCC	TTTCATTACT
177721	AGGGAGAGTA					
177781	TTCGGGCATG					
177841	TGGTGGGTCT	GGATCTTAGG	TAGATAAAGA	AACTTTAATT	CCACGATGTG	TTTTGGTAGG
177901	GATAGTTGGT	GGCAGGGATG	TCAGAGAGAC	TTTGAGGCTT	CTTCAGTTCA	ATATGACCAA
177961	GGGCCATATA	TTAGGGTATC	AATTTCTGAG	CCCCAACAAG	AGCTTAGGAG	AGATGTGATA
178021	GCATCACAGT					
178081	CCTGGCTGAA	GTACAATGGT	ACGATCACAG	CTCACTGTAA	TCTTGAACTG	GGTTCAAATG
178141	ATCCTCCCAT	CTAAGCATTT	CAAAGTGTTG	GGATTACAGG	CATGAGCCAC	GGTACCCAGC

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178201	CTGAAACTG	C ACCCACTTT	TGATAAACTT	TTCAAATGA	TABAGGGA	AGAGTAAGCA
178261	CIACICAGA	G GTAGGAAGAI	A AGGACACAGO	ATTATAGGA	יסממסממד ז	NCC2CC2222
178321	AAAACCAGA	C CGGTGTGGT	GCTCACACCT	GTAATCACAC	CACTTEGGG	GGCTGAGGTG
178381	GGGGGAGTC	A CTGGAGGCC	GGAGTTCGAG	ACCAGCCTGG	CCANCATACE	AAGACGCTGT
178441	CTCTATTAA	A AAAAAAAA	ACCTGCCTTC	AGCTAATCAC	CCARCATAGO CCARCATCAA	CCTGACAAAG
178501	GATGTCCCA	A AGTAAGTCTT	AGCATTTTT	, դերահանահանանարարում 1100111111111	CACACACHOMO	CCTGACAAAG
178561	CCCAGGCTG	A AGTTCAGTG	CGTGATCTCG	GCTCACTGC	ACACCECCE	CGCTGTGTTG
178621	AGCAATTCT	CCTGCCTTC	GCCTCCCAAG	TAGCTGGGAT	TACAGCIGCCI	CCCAGGCTCA
178681	CCTGGCTAA:	r TTTTGTTTT1	TTTAATAGAG	ATGGGGGTTT	CCCATCTTAL	CACCACCACG
178741	CTTGAACTC	TGACCTCAAG	TGATCTGCCC	ACCTTGGCCC	CTCCATGITAA	CTGGGATTAC
178801	AGGCGTGAG:	CACTGCACCC	GGCAAAGTCT	TAGCATTCT	TACARAGIG	TTGTACCCGT
178861	ATCTCTAAA	GGGAGTAGT	AATTTCACCC	CAAAATATGG	CTTCCTCTCT	TAATGAGTAT
178921	TTTGAATGA	AAACTCTTAG	AGATCAACAG	ACACTAAACA	CACTTOTIGATA	TAGGTACATA
178981	AAAATAGGAT	GGCCCACCA	GCGAGAACAA		CTCCCTCCC	TAGGTACATA GTTATCTCAT
179041	TGTGCATTAT	AGGAAAGACC	AAGAATGTAA	CCACACCTCA	ACACA CCCCC	TTATAAGATA
179101	ATCAGTCTCT	AAGCATCATT	TAAATTCCAA	GGAGAACTAT	TTACARCCCTT	TTATAAGATA ATCTGTTCTT
179161	TGATCCAATT	AGTCTCTCCT	GGTAGTTACA	TATTECCCCT	CARCAGATT	ATCTGTTCTT
179221	TGTTTCCCAT	AACCTATTT	GCAAGGATCA	AGCCCCCTCTT	CAACAGAATT	CCTCTTCTTC
179281	GCATATAAGC	TTCTAAATTC	CACTECEATA	TTCCTA CTA	ACTICITCAA	AGAACCACAG
179341	AGTAATTAAA	TTGTAAAGCC	מייייטייים מייייים מייייים	TENATETEC	GTGCATGAGG	AGAACCACAG TTCATTTTTC
179401	AGCAAAACTT	CCAAGGGCAA	AGGTATAAAA	CARACIGCC	TTTTTTTGTG	CCCCCAACCA
179461	TCTGAATAGA	CTTTCTCTTC	AGTCAGGCTT	CAAAAAAIAAA	ATTCTAAAGC	ACTGGCTCAG
179521	GCCATTAAGG	GAAGTGGGG	TTCAACATCC	CTIMAMATGT	AACCTGAAAG	ACTGGCTCAG TTAACATCAA
179581	CACAGCTTTT	AAGTCTGATA	ACD A A CATTOR	CICATTATTO	CTCTCTGGCA	TTAACATCAA
179641	TAAAAACTTC	ATCCCATAGT	ACA ACTOTOC	TACAACCTAT	TCTCTCTGAA	GCCTGCTAGC
179701	TCCTTTCTAT	TAATCCCAAA	TCTTTATACA	ARCTCACAGO	CTGTTATCAC	AACCTAGTGC
179761	ACTCCTCCGC	TGCTTCCAGT	TGTCCCGCCT	CTCTCCAACCA	ATTGTCATCA	CCTCCACCCC
179821	ACGTATTTGA	TTGATGTCCC	PECCECCE	A A A A HOME MA	AACCAGTGTA	CATTTCTTAA
179881	CACCTTGAGC	GCTTGTTCTC	AGGACCTCCCT	CACCCCCCCCC	AAGCCAAGGT	GCATCCCAAC
179941	AATTTGGCTC	AGAATAAATC	TCTTCAAATC	TTTTT	TCATGGGCCA	TGGTCACTCA
180001	AGATGACTGC	TTCACTGAAG	CCTCCTCTCC	A A CTICA CTICA	TITGGCTCTT	GTCATGACAC
180061	TCCCCGGATA	GCCCCAGAAG	CACCTACTA	TARTACACEE	GGGTTTTGCA	AGGATAATTT
180121	GAACACTTGT	TTTGTGCCAG	ACCURTORCE	ACATIMICACII	AAAGGTAGCT	AAAATGCATT
180181	ACTCCTGATT	TGTTAATACA	TTCTAAATAA	ACATTIGCTT	TGTGCCAGGC	TTATGCCAGT
180241	AAAACAGAAA	АТАААТАААА	ATATATATA	AAATTCTGGA	GTTTCAAATA	TAATAACTGA
180301	TGGTGGCTCA	CTCACACCTG	TARTATAGE	ACTGAAATAA	AAATTTACTA	AGGCTGGGGA
180361	CCAAGAGAGG	GTTCTTGGAT	CTCACACAAC	ACCGGAAAGG	GGTCCGTCCA	GATCCAGACC
180421	GCAAGTTTAT	TAAGAAAGTA	GAGGAATAAA	AAAGAATTUG	GGCGAGTCTG	TAAAGTGAAA
180481	TGAGGGCTGC	TGGTCGCCCA	TTTTTATGGT	TATTTCTTCA	TTATCTCCT	AGAGCAGCTC
180541	GGATAATTCA	TGCCTCCATT	TTTTAGACCA	TATABACTAR	CTTCCTCACC	AACAAGGGGT
180601	ATTCGTAAAC	TGTCGTGGCG	CTGGTATGAG	CATAGCACTC	ACCACCACG	TIGCCATGGC
180661	TCATCGCCAT	CTTGGATTTG	GTGGGGAGCA	CATAGCAGIG	AGGACGACCA	GAGGTCACTC
180721	CATCTTGGAT	TTGGTGGGGT	TTAGCCAGCT	TOTOMOGAIGA	TTTCCTTTT	CTCTCATCGC
180781	TTTTTTTTT	GCCCAGGCTG	GAGTGCAGTG	CCACCATCTC	ACCECATET	TTTTTTTTT
180841	TTCTGAGTTC	AAGCGATTCT	CGTGCCTCAG	CCTCCCAACT	AGCTCACTGA	AACCTCCAAT
180901	GCCACCACAC	CCAGCTAATT	ייייייערעינערעיייי	TTAATACACA	AGC I GGGATT	ACAGGCATGT
180961	TACGCTGATC	TCCAACTCCT	GCGCTCAAGC	CATCCAGGGA	CCTTACCOM	CCATGTTGCC
181021	GGGCTTATAG	GTGTGAGCCA	CCCCACCTGG	CCTAGCCA	TTCTTAGCCTC (	CLAAAGTGCT
181081	TATCAGCAAG	GTCTTTATGA	CCTGTATTTT	GTGCCCGGC	COTCOCTORS	CAACCTGTTT
181141	ACAATGCCTA	ACTTACAGGG	AATGCAGCCC	AGCAGGACTC	ACCOMPANDO (	CACCAACCAC
181201	CTATTCAAGA	TGGAGTCTTT	CTTGTTCAAA	TACCTCTCAC	ANGCCCARCE :	CACCCAGCTC
181261	ATGACACAGG	AGGATTGCTT	TAGCCTAGGA	GCTCABGACC	ACCCTCCCCC	LITIGGGAGG
181321	ACCCCATCTC	TAAAAAAAA	AAATACAAAA	ADATTAGECE .	RGCATGATCC :	ACACAGTGAG
181381	TAGTCCCTGC	TACTCAGGAG	GCTGAAGTGG	GAAGATAGCCA	TCAGCCCACC :	LGIGIGCCTG
					LUNGCCCAGG A	AATTCAAGGC

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181441		GAGGCATTTG				
181501						CACAGGATGA
181561		CAGCACAAGG				
181621		GCCAAAACCC				
181681		ATTATATGTT				
181741		GCTTACAAAT				
181801		CCCTCAATTT				
181861		GTTTAGCACA				
181921		TTCTGCCTAC				
181981		TTTACTGTAT				
182041	CCTCTCTTGG	GGTCTGGATC	AAGACCCCTT	TCTGGTAACA	TCTTTCTGGT	GACCACGAAG
182101	GGACAATACT	GAGGAGACTC	TGAAGCCAAA	GGAAACAGAC	TACAGCACCA	ACTGGCTGAC
182161	TTTGGGTAAG	TGGTGGAGTC	CCCGGGTAAA	GGATAGGATT	GGGTTAGAGG	TGCAACTTAG
182221	GGGAGATAGG	GTCTCTCCTA	AGACAGAGAG	CGTTTCAGTC	CGCTCTTAAT	AAAGGGCAAG
182281	AATGCTTGAC	CGAACTTGGG	TTTGAGACCC	AACTTAGGAA	GGCTACAGTC	CTTAAGATTT
182341	AAGGGGTTAG	AGGCCCCTCT	CAGTAAAGTC	TCTCTTGGTT	AAAAACGGAT	TTAGCATTAG
182401	GGGATGTTAA	CTGCTATTCT	GTTTGTATTA	ATCTTCCCTG	TGCTCTTTGC	TGACAGCTAT
182461		ATTAGGCATG				
182521		TTGACAGCTG				
182581		TTTTGATTCA				
182641		TTCCCCACCT				
182701		TTCTGTTACT				
182761		GTTTGATTAA				
182821		TGGGTGCTAA				
182881		AATGGAAAAC				
182941		GAGACATTTA				
183001		TAGCTTGGAC				
183061		ACCCAGAAAC				
183121		CTCTCTCTCT				
183181		GTTTTGATTA				
183241		TACTTTGTGA				
183301		ATAGAACATG				
183361		CGTTGCAAAG				
183421		CCTTCTCATC				
183481		CTCTTGGTCT				
183541		AAAATTATCT				
183601		GCAATGGAAA				
183661		GGCCAAGGTT				
183721		TTACCATTTG				
183781		TGAGAACCTG				
183841		ATGGCTAAAG				
183901	CTTTAGAGTG	CTATGGTTTA	TGGTTAAAAG	ממדדממדדים	AGTGGATATT	CANTCTCTAA
183961	AAGCCTGGGA	CTCCTTGGGA	AAAGCAGAGG	AGGCACCACA	GACCCCATTT	TOGGANANCO
184021		TCATGAAACC				
184081	GCTCTGTTTG	GCTTTGCATT	ATGTTATCTG	ארכיייייייייייייכא	CHITTERGGGG	TATCACAAAT
184141		TATGAGGGAG				
184201	GATAGCTAAA	GGCAAATATA	GGTGAATACT	ተርርርጣን ተተተ	CITOCATALTY	TCACARCARC
184261	CATTCTCTTG	ACTACCTAGA	AGGTATGGAN	TOGCINIIIG	CCCCACCGAC	ACATAACATT
184321	CCCAGGGGAG	ATGGCTGATC	CCCCAAAAGA	GGGCTGATTC		CATCCACCAT
184381	CTGGTATAAA	AATGGGACCC	TGGCCAGGCA	CAGTGGCTGATIC	CCICILIIGG	CTCDACAGGAI
184441	TGGGAAGCCT	CAGAGTTATG	AATGTCTCAC	CATACTCACA	CACCIGIANI	CICAACACII
184501		CACAAGAGAC				
184561	TGAAGAAGTT	ATAGAAGATG	GATCTTTATC	CCACTGCAAT	CCTTACCA	ARGGETTEEE
184621	TGGTAAAAGG	GAGTGGGAAA	ATATGTCAGA	GGCATTTCAN	TCAGAGTGAC	TCCATCTTCA
						* ~ ~~ * * * * * * * * * * * * * * * *

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184681	NTN GGGGGG					
184741	ATAGGGGCT	G GGTAAAATAI	A GGCTGAGGC	C TGCTGGGTT.	A GGTTAGGCA	TCTAACCAGG
184801	AGITIAGIC	A CAGGATGAG	A TAGAAGGTT	G CACAAGGTA	C CCCTCACAA	Caccamacama
184861	AIAAAAIAG	G TAACGGTAA	A GAAGCCAGC	T AAAGCCCAC	ר אאאסרכאארי	TOCOCOONS
	AGIGACCIC:	1 TGTCATCCTC	: ACTGCTCAT	A TACACTAAT	ד אייא כידיכי היייו	ACCATICOTAC
184921 184981	MAGACACTC	CACCAGTGC	: ACGACAGTT	I ACAAATACC	A TGACAACATC	TOCACOTTAC
	CITATATEG	r CTAAAACGGG	GAAGAACCC'	TAGTTCTGG	S AATTCTCCXC	CTCTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT
185041	MANAGE TO T	GAATAATCC	I TTAGTTTAG	C ACATAATCC	מתיים ממדמת במ	TROCHOROGE
185101	IMITIGAGCA	4 GICCATACTO	CTGCTCTGC	TATGGAGTA	יייייייייייייייייייייייייייייייייייייי	
185161	IMITITIAL	S ATAAAGACTO	GCTCTGTCA	TCAGGCTGG	GTCTCCACTC	Chaman dama
185221	1111000101	A CIGCAACCII	CACCTCCCG	GTTCAAGCA	THUTCHICA	TO COMPAGE
185281	WC TWOC LOC	F GACCACAGG1	GGGTGCCAC		א ערייטיעיעיעיעעע ע	TITLE THE SAME OF
185341	MGM I GGGG I.I	TCGCCATGTT	' GGCCAGGCTC	GTCTCGAACT	CCTGGCCTCA	3.CCC3.MCC3.C
185401	1100011000	CICCCAAAGT	GCTAAGATT <i>i</i>	\ CAGGCATTA(	י לרבריים יוכרים	TC 3 CC C3 mm c
185461	TITIMITICI	TAACTTTTT	TTGTTTTT	GAGACAGAG1	بالمالية ليناسك لأماليان	C3.0003.000
185521	AGAGGC I GGA	I GIGCAGIGGI	GCGATCTTGG	TTCACTGCAR	CCTCTGCCTC	CTCCCTTC
185581	GCGATICTIC	. IGCCTCAGTC	TCCTGAGGAG	CTGGGACTAC	LAGACATOTOC	CICMINGIAGO
185641	AGCIAATITI	GTATTTTAG	TAGAGACAGT	GTCTTGCCAT	GTTTCTCACC	CTTCTCTCTCC
185701	ACT CCT WACC	TCAAGTGGTC	TGCCTGCCTC	: AGCCTCCCAA	ACTCCTCTCT	TTT CT CCCT
185761	MAMICACIGO	GCTCGGCCCT	TCTTTACTT	'CTTAATAAAC	ר ערות התהתהות אירי	TOTAL COLORS OF
185821	GGACTAGCCC	CAAATTCCTT	CTTGTGTGAG	TTCCAATAAC	ملتحاملات المتملميات	CMCARAGORA
185881	TIMIGGEIGE	TGTTCAGGCT	GGAGCAAGCT	'GGAGCTCATG	CTCCTCCTCA	CACCCCACCA
185941	racarakici	GIGATCCCAG	TAAGAGGATC	ATGGTCACTC	CAGCCTGAAC	Chancamas
186001	TAICICATCT	GTAAGAAAAA	AAAAATTACT	' AGAGGGCTTT	AACAGCAAAT	TTCACCACCA
186061	AMMAGAAGTA	ATCAGTGAAC	TCAAAGATAG	GTCAATTGAA	ATGATOTACT	CTCABAAACA
186121	GAAAGAAGAC	AGAATGAAGA	AAAAGAAATA	GAGCCTTAGA	GACAGGGGAT	ACCAMONACA
186181	MINCIANTAT	ATGCATAATG	GGACTCCTAG	AAGGAGAAAA	GTGDGDGGDC	ACCCRORGE
186241	AATGTTTGGA	GAAATAATTT	CTCAAAGCTT	CCCATGTTTG	GCADANANC	AGGGAGAGAG
186301	ATACATATTT	TAGGAGCTCA	ATGAATTCCA	AGTAGGATAC	ACTCARAGAC	ATTAACTTGC
186361	AGACACATCA	TAATCAGATT	ATCAAAAGAT	GAAGAAGATG	AATCTTCACA	CCACATACCT
186421	AGGAACAATT	CATCACATAC	AAATAGTACT	CAAAAGATGT	CTGGAGTAGG	TATA CTA ATA
186481	TCAGACAAAA	TAAACTTTAA	GATAAGCATT	GTTATAATAA	ATABAGAAAG	CTATTCTAATA
186541	ATGATAAAAG	TGTCAATTCA	TCAAGAAAAC	ATAACATTAT	AAACATACAT	CCACCORAGO
186601	ACAGAGCCCT.	AATATTCATG	AAACAAAACT	GACAGAATTG	AACCCACAAA	TACASABOOG
186661	GACAATAATA	GTTGGAGACA	TCAATACCTC	ACTAGTTAGA	CAACATCAAC	IAGAAAATTC
186721	AAGACTTAAC	ACTTGAAAAC	ACCTAACCTG	ACCCTAACAT	ANTONICANC	AAAAAAATAG
186781	CCCAAAACAG	CAGAATAAAC	ATCCTTCTGA	AGCTCACATG	VVVCVUMMAN	GICACTACAC
186841	TGTATATTAC	TTCATGAAAT	AAGTCTCAAT	AAATGTAAAA	CCACRAMA	CAGGATAGAC
186901	ATATTCTCTG	ACCAAAGTGG	AATGAAGATA	CADATCAATA	ACMA GGOGG	AATAGAGTAT
186961	TCACGCCTGT	AATCCCAGCA	CTTTGGGAGG	CCAAGGCGGA	CACAMGACTGG	GCGTGATGGC
187021	TTGAGACCAG	CCTGACCAAC	ATGGTGAAAC	CCTGTCTCTA	CAGAICACGA	GGTCAGGAGT
187081	GCCAGGCCTG	GTGGCATCTG	CCTGTAGTCC	CAGCTACTCG	CIAACAAAAT	ACAAAAATTA
187141	CACTTGAACC	CAGGAGGCAG	AGATTGCAGT	GAGCTGAGAT	CCCCCCACTGAG	GCAGGAGAAT
187201	TGGGAGACAG	AGCGAGACTC	CATCTCAAAA	TTANANAAA	ARRICACTO	CATTCCAGCC
187261	GAACAAATCA	AACCCAAAGC	AAGCAAGAGG	ANANTONNA	AMAMGAAACT	AGAAAATAA
187321	AAAAGGCACA	TTATGTACAG	AAGAACAAGT	CTATACATCA	ATTTCAAAGC	AGCCAAGAAC
187381	TATAAGCAAA	AAGACAGTGG	AGCAAAATTT	TTTTTTTTTTTTTTTT	CATATTTCTC	ATAGACACAA
187441	ACCAAGCAAA	AAAACTCCCC	CCADATGAGG	CTCARATIAA	TGAAAGACCT	ACAATTCTGT
187501	GAGGAAGGAA	TTTATCTAGT	CATATETEAG	PUTTONALINAG	ACAATTTAAT .	ACAGAGAAAA
187561	TGGATGTTTT	CTATTTCATT	TABLETALL	VCCCCCCS zw	TACATTTTGT .	ACTGTATATG
187621	TTCTTTTTGA	TTGACACAGT	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	ACCOLUCAAT	AAATGGTAG	ATTGTCTTGC
187681	TAAAGGCAAT	AAACATCTAA	TCAGCAGACT	ACIACIAG	ATTTTTTA '	TCTCCCTGCC
187741	TTTAGGCAGA	ATGATAAAAC	TCCCTTA CCC	AGMACAATAA	AAAATATTTT '	ITAAAAGTCC
187801	AAACAGTACT	AGAAATTGTA	TOUTINGGC	TABACACAM	TCCTATTTAT A	ACAAAGGAAT
187861	GTGGTTGACT	ATTTTCACAN	A KUMUNATAA	AMMACAGATA .	ATATTTTTC 7	CCATAAAAT
			ningiinh	CARIGIARIG	IGTGATTTAT !	AGCATTTAAA

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187921	AGTAAAACAG	GCCGGGCACA	AAGGTTCGTG	CCTGTAATCC	CAGCACTTTT	GGAGGCCGAG
187981	GCGTGCAGAT	CACTTGAGGA	CAGGAGTTCA	AGACCAGCCT	GGCTAACATG	GCAAAACCCC
188041						AATCCCAGCT
188101				GAATCCAGGA		
188161				AACAGAGCTA		
188221				CAGTGCAAAA		
188281				TGAACTACGA		
188341				AACGAAGAAT		
188401				TAAGTTGTTC		
188461				AATGGGAAAG		
188521						CTAATACAAA
188581				CAGACAACAA		
188641				TTTAATATAA		
188701				CTCCTAACTG		
188761				GACAGAGTCT		
188821				GCCTCTACCT		
188881				ATAGCACATG		
188941				ATGTAGGCCA		
189001				GCGTGAGCCA		
189061				TTTGTGCATT		
189121				GTCACTGAGG		
189181				AGCTGGGGTA		
189241						
189301				TTTTTTTTT		
189361				GAACATGGCT		
189421				CTGTATAGCT		
189481				TTTAGAGATG		
189541				CCTCCCACCT		
189601				AGACTTTTTA		
				GGATTGATGA		
189661				TGCTCTGCTG		
189721				TTCTAGTTTA		
189781				CTTATTAGAC		
189841				TACTATAAGC		
189901				TTTTGATGTG		
189961				TTGTTTGGAG		
190021				CCTACATTTC		
190081				TCATTTATTT		
190141	GAGACAGGGC	CITTCTGTGT	TGCCCAGGTT	TGTCCCAAAC	TCCTAGTCCC	AAGCAGTTCT
190201				ATAGGCACGA		
190261				AACAGACATT		
190321	TTGGCCAACA	AATATATGAA	AGAATGCTCA	ACATCACTGT	ATTAGTCTGT	TTTCATGCTG
190381				TTTACAAGAG		
190441				TCATGGTGGA		
190501				AGAGCTTGTG		
190561				TATCATAAGA		
190621				CTCCCAGGAC		
190681				CAGCCAAACC		
190741	GGGAAATGCA	AATCAAAACC	ACAATAAGGT	ATCATCTCAC	CCCAGTTAGA	ATGGCTATTG
190801				GAGGATGTAC		
190861				CATTATGCAA		
190921				AAAGGAAACA		
190981	GAGAGAACAA	ATCTCTTGAC	ATTACACAAA	CTGCATCTGG	GGCTAGTGGT	TAGAATATCC
191041	TCAGTCAAGG	AGGTAGAAGA	GCAGGAGGGA	AAATCCCTAA	GTTCGTGCAA	GTGCAGAAAC
191101	CCACAAGCTG	TGTTCTCAGG	TTGACATATA	CTCATTTTAA	TAGTAAGAAA	CACACCCTTG

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191161	GGTAGAGAAT	TAAAATGCTA	ATAATACATG	TGATGTATGT	ACTAGCGTGT	ATGGCAATAT
191221	TGCATGCACA	TTCAAGAGAC	CACCCAAAAC	ATATTTAACA	ACAATGCCCA	TTCCCACCCC
191281	CTCATGGATA	ATCACGTAGG	ACTCCCATAA	CGGGAGTTTC	TTCAGTGTCA	ATTGGTGCTG
191341	AAGTAGCCGA	CCCTGACTCT	GCTATCAGCG	TGTACTTTCA	CCTTGCAATA	AACTCCTTTG
191401	CCTACTTTTA	CTTTGGACTG	GCTTTCAAAT	TCTTTTGTGC	AGGGAATTCA	AGAATCTGAA
191461	CCAGCCCACT	GACAACAGAG	GTTTCTCAGA	AACCTAAAAA	TAGATCTACC	AGATGAGGCT
191521	GAAAATCTGC	TACTGGCTAT	TTATCCAAAG	GGAAGGAAAT	CAGTATACAA	AGAGACACCT
191581	ACATCCCCAT	GTTTATTGCG	TCACTCTTCA	CAAGAGCTGA	TATATAGAGT	CAACCCTAAA
191641	TGTTCATTAA	CAGACAAATG	GATAGAAAAT	GTGGCATATA	TACACAATGA	AATACTATTT
191701	GGCCATGAGA	AGAATGCAAT	CTTGTCATTT	GTGGCAACGT	AGATGAAACT	GGAGAACATT
191761	ATGTTAAGTA	AGATAAGCTA	GGATTGGAAA	GATAAATACT	ACATGTTATC	ACTCATATGT
191821	GAAAGTAGAG	AAAAATTTTT	AGCTCATGGA	TTTAGAGAAC	AGAACTGTGG	GTACCGGAAG
191881	CTGGGAAGGG	TAGCAAGGAG	GGGAGGATAG	GGAGAGGTTG	GTTAATGGTG	ACAAAATTAC
191941	AGCTAGATTG	TAGAAATGAG	TTCCGGTGTT	CTGCACCATT	GTAGGGTGCA	TATGGTTAAC
192001	TCTCATTTAT	TGTATATTTT	CAAAAAGCTA	GAAAAGAATT	TTGAATACTC	ACAACAAAAT
192061	AAATGATAAA	TGTTTAAGGT	GATGGATATA	CTAATTACTC	TGATTTGATT	ATTACACATT
192121	GTGTACACAT	ATAAAAATAT	CACTCTTTAT	CCCGTATATA	TGTACAGTTA	TTATATGTCA
192181	ACTAAAAATA	AAAGAAAAA	AGAATATGAT	CTATCATGAT	GTATATATCA	TGTGTACTTG
192241	AGCAAAATGT	GCATGCAGAT	ATTGTGTATA	ATGTTCTATA	AATCAATTAG	CTCAAGATAA
192301	TAGATAGGAT	TGTTCAGATC	TTCTGTGTCT	TTACTGATAT	TTTGTCTAGT	TATTGCATCA
192361	TTACCAAAAA	AAGGGTGTTA	AACTCTCCAA	ATGTGATTGT	AGAATTGTCT	ATTTTGTCTT
192421	TTCTTTTCCA	TTTTTACTTT	ATGTATTTTG	AAACTCTGTT	ATGACATTTT	GCTATGTATT
192481					TCATACATTT	
192541	TGTTTTCTTG	ATGAAATGAC	CCTTTTCTAT	TGTCGTTGTT	TTTGTTTTTT	CTGAAATGGA
192601	GTCTCACTCT	GTTGCCCAGG	CTGGAGTACA	GTGGCACAAT	CTTGGTTCAC	TGCAACCTCC
192661	ACCTCCTGGG	TTCAAGCGAG	TCTCCTGACT	CAGCCTCCAA	GTAGCTGGGA	TTACAGGCAT
192721	GTGCCAGCAT	GCCAAACTAA	TTTTGTATTT	TTATTAGAGA	CAGAGTTTCA	CCACGTTGGC
192781	CAGGCTGGTC	TCGAACCTCT	GACCTCAGGT	GATCCGCCCA	CCTCGGCATT	TTTATTTTAT
192841	TTTATTTTT	TGAGACAGAG	TCTCACTCTG	TCACCCAGGG	TAGAATGCGG	TGGTGTGATC
192901	TTGGCTCACT	GCAACCTCCG	CCTCCTGGGT	TCAAGCAATT	CCCATGCCTC	AGCCTCCCGA
192961	GTAGCTGGGA	TTACAGGCAC	ATGCCACCAT	GACTGGCTAA	TTTTTGTATT	TTTAGTAGAG
193021					TTTAACAATA	
193081	CTCTGTCTCT	GGTAACACTC	TCTGTCTTAA	ACTCTATTTT	AGCTGTTATT	ATTATAGCCA
193141	TTTTAGTCTT	TTTATGCTTT	CTGTTTGCAT	AGTGTATATA	TTTTAATATG	TTTATTCTCA
193201	AGTTATCTGT	GTTTTTATAT	TTAAGATGTT	TCTCTTCTAG	CCAACGTGTT	TGGTTCTTGC
193261	ATTTTTAAGT	CGATTCTAAC	AATCTTTGCC	TTTCAATTGA	AATATTTACA	CCATTAACAT
193321	CTAACATTAA	CATTTATTTT	TCTTTCCACA	GTACACTGGC	TAGCATCTCC	CATATAATAT
193381	TGAACATAAA	GTGTGATAAC	TGACATCCTT	ATTTCATTCC	TACTCTGAGT	GGAAAGGGCA
193441	GGGGTGGAGA	AAGCATTCAA	CAATTTGCCA	TAATTATAAT	TCTTTTTGTT	ACACTGTTTT
193501	CTTCTGCATT	AAAAAATATC	ATTACATTTT	GCATGAATTA	TTAGGAGAAA	ATATTTTCCA
193561	ATTTTCCTGG	AAAATGCCAT	AACCACGTCT	CTCAATTTTG	TTTCCATCTT	TCTTCCACAT
193621	TTTACATAAC	CTACATAAGA	GACACATTAT	CAAGTATATT	TTACATGGCT	TCTCAGTGTC
193681	TTCTCTGTCT	GCTAACAGGT	TTACCAAGAG	ATGGCACTCT	TGTATTTCTG	GTGGCTATGT
193741	CCATATCGTT	TTGCCTTTAA	GACAGCGTAA	CTACTTCTTT	CACCAGTATT	AAAGACATGT
193801	ACATTTGATC	TGGTTCTTGT	GGATGATTTT	AAATGACTCA	AGCTAATAAT	CCTAATTTTA
193861	CCTAAACACT	CCATTATTTT	AAAATGTATT	CCTTTATGCC	CACAATAAAC	ATTTATTGAC
193921	ATTAGGCTGG	ACATTAGGCT	TCTCTATGGC	AGACATTAGG	CTGGACCCTA	GCCATATATC
193981	TATTGAGGGA	AAAAAAATTA	TTTTCTATAT	AAGTTTCCAG	AAAGCCAAGA	TGTGTTTTAA
194041	AAACAAAACA	AAACATTACA	TTCTAAATGC	TGTAACAAGA	TAAGAAAAAG	TGTTGAGGCT
194101	GAGAGAAGAA	CAAAGCAGCA	AGCAACTCCT	GGAAGGACCA	CTGCTGCAGA	GGTAATAACT
194161	GGTGAACCAT	GITTTGGAGA	AGGAAAAGGT	CACCAAGAGA	AGGAGGGGGT	CCAGGGTGTT
194221	CAGAAAGATT	GCATGCATAA	AGATCAAGGG	AAAAAATAAT	AATTCCGTAT	TATGTAAATG
194281	TGAAGTTCCA	GGACCATGAG	CTTGGAGAGC	ATGAAGTACA	GGAGGAGGGT	TGGTTTCAAA
194341	TAAATCTGGG	AATGAAACAG	TGAAGCCTCT	GGCAGAACTC	ACATCTCTTT	CCTCCCCTCT

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194401	TCCTTGCAC	A TTCCCTTTAT	GGAGTAATTO	CAGGGATGGG	AAAAGTTCAA	AACCACCACT
194461	GAGCCTAGG	A AGTGCTAGGG	TAAAGTGGAG	AATGAACCTG	CGTGATTTGC	TCATCCTAAA
194521	CTAGGTTCTT	r ctaggagagc	CCTTCCCCAI	AAAATCTGCC	CTCCTCGAAG	GGGCCCNGAC
194581	AGCCTAAGCT	r cacciccaa	AGACCCCTTA	CTTGCTGACT	GAATCTGATT	CCACCCAGAC
194641	ATGGCCTAA	A ACCCTTCCAT	' AACTCTATAG	CCAAATTCAA	TTTTAGACAG	GCCTCATACC
194701	AACCTTTCTT	CCTCTAAGTC	TGCCACCCTA	GGCAATTCTC	AACATTCTCT	ACACACTTTC
194761	GGGCCATAGA	CGTGCTACCA	AGTCTCCAGA	CCTAGACCTG	ATGGAGCAGT	COTOTANTON
194821	GACGACCACT	GGCCTTTGAA	CCAGACCCTT	CTCTGTGGCT	CCTATGCATC	TOCARCOTOR
194881	TTTGAGCACT	GCTGCCAAGA	CATCTTTGGC	ACTITGTTGT	GARGTTTTAN	A A CTCA A CTA
194941	ATCTACAAAA	CACCTAACCT	TTAAAAATTC	ATTGTCATTT	CATATCATCA	DAGATARACIA
195001	AAGGCCAGGA	AACTGTTCCA	GGTTAATAGA	GACTAAAGAG	ATAGCAACCA	ANGATAAAGA
195061	GTGATCCTGG	ATTGAGGGGA	AAAAGTGTTG	TCAGAGACAT	GATTGGGACA	GCTGGTAAAA
195121	TTTGAATTTG	AATTTAAAGA	TAAAGTATTG	AGTAATATAG	GAAGATGATT	ATCTCCAACT
195181	TTCAAATGTT	TCAGTAAGTA	TATATATATA	TAAAGAGATA	TAAAGACATA	TANATANATA
195241	GATGGATAGG	TAGAGAAAA	GCAAATGTAT	AATATTAACA	ATCTAGGTAA	TANGETHANIA
195301	AGTGTTCTTT	GTACTGTTTT	TCTGATTTTT	CTATATGTTT	GDADTCATT	TARAGIAIAIG
195361	AGGTTTTTGG	GGTTTTTTTG	TITGTTTTTT	GTTTTTAGAG	ACACCATOTT	AMMATAAGA
195421	CCAGGCTGTA	GCTCAGTGGC	CCAATCATTG	CTCACTGCAG	CCTCXXCTTC	ATTCTGTCAC
195481	GTAATTCCCC	CTACCTCAGG	CTCATGAGTA	GCTGGTACTT	CAGGTGTCCA	CIGGGCTCCA
195541	CAGCTAATTT	TTATTTTTA	AATTTTTGTA	GAGATGGCAT	GTTGCTATCT	CACTGCACT
195601	AGTCTCAAAC	TCCTGCCCCC	AAGTGATCCT	CCCACTTTGG	CCTCCCAAAC	TOTTACAATT
195661	ATAGGCATGA	GCCACTGCAC	CCAGCCCCAA	ATANANANGT	Dummer Valenting	A A TOTA A CORN A"
195721	TTAATTTTGA	GTCAGAGTTT	CACCCTTGTC	ACCCAGGCTG	GAGTGCAATG	CONTONTONT
195781	GGCTCACTGC	AAACTCTGCC	TCCTGTGTTT	AAGCGATTCT	CTTGCCTCAG	ACTCCTCACT
195841	AGCTGAGATT	ACAGGTGCCT	GCCACCATGC	CCAGCTAATT	יייייייייייייייייייייייייייייייייייייי	TACTACACAC
195901	GGGGTTTCAG	CATGTTGGTC	AAGCTTGTCT	CAAACTCCTG	ACCTCACCTC	ATCCACCACA
195961	CTCGGCCTCC	GAAAGTGTTG	ATGAGCCACC	ACACCCGGTC	TADADACTAT	TTTDDDDDDDD
196021	CAGTCCCACT	CTACCTTGTC	CTACACTACC	AGGGGCTAGG	ATCACCCCAT	CTCTTCTACC
196081	CTATGAGATA	GAGGAATCCA	AGGAAGAAGA	TAAGCTACTT	GCTTCCTCTA	TACCCTCTTC
196141	TGTGTGCTCT	CATGTGCTCT	CTCTCTCTCT	CTCTCTCTCA	CACACACACA	CACACACACA
196201	CACACACACA	CACACACATG	AATACCAGAG	CTATCACTTT	CCCAGTCTAG	TACTCATACTC
196261	ATCCCAAGGG	TTTTGTGTTG	TAGTGGTTTG	CTCATTTGTT	TGTTTTGTTT	GTTTGCTTCC
196321	ATTATTCTTT	TTCTCTTTTT	GCAGCTGAAG	GGAGAATTTC	CAGGCCAGCC	CTTTCCCCOAT
196381	TAGAGTTACA	GTGCCTCTAT	TCAGGCTTCA	TAGAGAGACC	TEGENTTONE	TACTCCCCCC
196441	CTTTTATCCA	GTTCAAAATA	ATGCATTCTC	ACCAAGATGT	ACTITICADAT	AAAAAAAAAA
196501	TAAAACACAA	AATTTTATTT	ATGCTGAACA	TTGAATCACT	TTTTTTTTTT	TTTTTCTCTTC
196561	AAAGTTATAC	ACACACAAAC	ACATTTGCTC	CTGCTTTGTT	TATTGGCCCA	CCCCTATCTA
196621	TGGTAATACT	TCATCAGGCA	TGAGTAGTAC	GTCTTGGAAG	GTGTGGTCTA	AAGCCTAGAC
196681	TCCTATCTGC	TTCCTTCAGC	ATTCTCCAGT	GTATCTGTCA	TCTGTCTACC	TTACCATCC
196741	GTCTCCAGAA	CTTCCATTCA	CATTTAGAAG	AGGGCAGCGG	CTTTCTATGG	AAAATATGAA
196801	CTCTCATTCA	TCTCTATTCC	TTCTTCTAGC	TATGGTCCAG	CTCAGCTGTT	TGGDATAAAG
196861	TATCTATATG	AAGTCTGCGA	ATGGTTCTCA	GACTGGTTGA	ACATTAGAAT	CACCTCACTA
196921	CCTTCTAAAA	TTCTTATTAC	CCAGGGCATA	TCTCAGAATG	AGTACCACAG	CCTACCCATA
196981	GGATTAGGGA	TCATGATCTC	TGGAGTCTGG	TTTAGGCACT	AGTGCTGTTT	ADDODAIA
197041	TCATGAGGTG	GAGGTTGCAG	TGAGCCGAGA	TGGCGCCACT	GCACTCCAAC	CTCCCCCACA
197101	GAGTGAGAGT	CTGTCTCAAC	AACACAAAAC	AAAAAAAACC	AACTACCCTT	CIGGGCGACA
197161	TGTCCATCCA	AAATTGAGAA	CCATTAGGTA	AGGCCAAGCT	GTATAATTAA	AGAGCAGTTT
197221	TCATTTGTCT	GGTGTGGTGG	CAGCTTTTTG	ATAAGGGAAG	TATTGTTGCC	TOUCHOIII
197281	CTGAGCCTCA	CTCCTGAGAA	CACTGGTGTG	TATGTTGCTA	AAATTCCCCA	ここかにとかかいける
197341	AGGITCCTTC	CTGGATAAAA	ACCACTGACC	CTGGGAATGT .	ACCCACTGCC	AATCTCCTGC
197401	GTAAACCTTG	GATACTGGGA	AGCCTACAGT	TGAAAATATT	GGGCTTGAGA '	TCCTGAAACA
197461	AATCTTGTAT	TTCATTAAGA	CTAATATTTG	GTACAGTGCA (	GCAAATCAAG	GAATTTTGG
197521	TGGCTGAGTT	CTTTTAGAAC	TTTTGCATTG	AAATAGGTTC	AAGCAGCAAT	DOGTTANANC
197581	TACAACCTCA	GCTAAAGGAT	TAAAAGACAC	GTGAGCTGGG	TAGGATGAGG	TOTAL LANGE
		_				CINCOLLIA

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197641	GGTGTGGCG	CTCATACCTG	TAATCCCAGC	ACTTTGGGA	ACTGAGGTGG	GTGGATCACT
197701	TGAGGTCAG	AGTTCAAAAC	CAGCCTGGCC	AACATGGTG	ADACCCATC	CTACTAAGAA
197761	TACAAAAAA	TTAGCTGGGC	GAGGTGCCAG	GCACCTGTA	TCCCAGCTAC	TGGGGAGGCT
197821	GAGGGAGGAC	AATCACTTGA	ACTCAGGAGG	CAGAGGTTG	r recendence	GATCGCACCA
197881	CTGCACTCC	GCCTGGGTGA	CAGAGCAAGA	CTCCATTTAL	AGIGAGCIGA	ATAATAATAA
197941	CAATAATAAT	AATTCAGACA	TATCCAGGCA	TCADACACA	NAMMATAATA	GATGAATAGT
198001	CTTGAGATTC	AAGTCACACA	TGAAATTTAG	GTGGAAAAT	ACCIDEDANA	AATTTGAGAT
198061	TATGATGAAT	GGAAATTTTT	CAAAGAGGAA	TTTCAGGCT	TOTTOTTOT	GGGATAGATG
198121	GACTTCCAAC	AGCAATAACA	CAGGATTAAT	GAGGACTTC	CATCTTGAG	GGGATAGATG AAATTAGAGA
198181	TGTTAGATGG	ATAAAGAGAT	AAAAGTACTC	TOTOTA ACA	CATGGGACAT	GAGATAGAGA
198241	CACTTCTAAC	CATCAGATAT	AACTAGCAGA	CTANACCCTC	CAIGGGACCA	AATCATGCCC
198301	CACTCCTGCT	TAAGACATTT	TAATTACTCT	CAGTAACGGIC	TAAAAATAAA	TACTGTGTTA
198361	TCTTTAACTA	CAGGGTTGGT	CTGGGTGTGC	DACACARCA	1CAGTTTTTC	TACTGTGTTA ATACATGGAT
198421	TCAAGTGTAT	GCCATGTACA	CCTATTCTT	AACACACAAAA TATOTTOTOTO	AGCCTGGCAT	ATACATGGAT CTTTTTCACA
198481	TCTGTTTTT	CCTTCATTCA	DGTCNDTCCC	TCATGIACIAI	TICATGIATT	CTTTTTCACA
198541	AGTTATATAT	AATTGTTACA	ABACAAAGGC	CCARARAGA	TTCTACTATT	CATGTGTACT AGCAACACAC
198601	ATTTATTATT	ACCTAAGGTC	TOTOGRAPACE	ACTUAL CONTRACTOR	AGTGGCTTAA	AGCAACACAC
198661	CTTCAAGCCT	CATGTGGCTG	CARTCCACCT	AGTTCTGACA	TGGCTTAACT	GGGTTCCCTG
198721	TTGATTGTGG	CATGTGGCTG	TTCCAACCTC	GTTGGCTGAG	TCTGAATTCT	CATCAGAGGC
198781	GCACCGGTAG	AAATTTCCAC	CTTCAAGCIC	CCTCAGGTTT	GTTGAAAAAT	TCAGTTCTTT
198841	TGTCACCCAG	AAGCTTCTTG	BCTCC COCK	ATTCAACTTC	TAGAGGCTGT	CTGCAGTTCC
198901	AATCAATCTG	GGTGGAGTGC	TCACCATCCT	TCATAGCTCA	CTGCAGCCTT	GACCTCCCAG
198961	ACACCTGCCT	TTCTCCCACC	CARCCATCCT	GAGTAGCTGG	GACCACAAGT	GTGTGCCATC
199021	CTGGTCTGGA	AAAAAACAAA	CAAACGAAAA	AAAACCCCCA	GAGAACTTTG	TAGAGACAAG
199081	ATAGGTATAA	ACTCCTGCGC	CEGGGATT	CTCCTGCCTT	AGCCTAAAAG	TTCTGGGATT
199141	TTTATGTCTG	GCCACCATAC	CIGGCATATG	GCAAGTCTTG	AGCAGGACAA	ATACAGATGA
199201	TCCATCTATT	TCTTCCATGG	AATTCTAGGT	TATTGTTGAG	ATGGTCCTCT	ATTGTCTTGT
199261	GTGTCTCTCT	GATTAGATAA	AACGITGITC	CTTCTGTTAT	TTTTCAACAG	TAGCTTTTAT
199321	CTTTGGTTGA	TTATCTTAAA	CCTCTTCTTTCTTTC	AAGAGCTGCT	CTTTTCTTGG	TGTACTTTAC
199381	CAGATGTGAG	TCCTTCTTAA	ACCIDITETIC	CCCTCTGGGG	CCTAAGATGA	GGGCTGTTAT
199441	GTCTAGGTAG	TCTATGGGAA	TCCCCCTTCC	AGGITCTTCA	GCCTCCGTTC	AGCCTTAAAT
199501	GGGTCTCAGC	AAATCAGTCA	TCCCCTTTCC	AATGTGGTAC	AGACCAGATC	ACAGAGACAG
199561	GGAGAACTCC	CAAGGTCTTG	CTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	CTTATAGAAA	TAATGAGTGT	TTACTTACTT
199621	ATGTCTTGGG	CTTGGAATAT	TAGAGGGATT	GAACCTGAGG	CAACTTTTGG	TGATTTCTTG
199681	TTTTGTGACC	AATCTTGGTC	TAGAGCCATT	TCAACCTGAT	TTCTTTTCAT	GTCAGTGGCA
199741	TGATGTGAAG	AGATAGTAAA	TCACCCCCCC	GATGTTCACT	CAGAGAAATA	CAATGACTTA
199801	GTCTCCTGCT	CTTCTGTGGT	ACTOROGORA	CITCATCTTC	ATTCCCTCTT	ATCTGCATCT
199861	GTAGCACTTA	TGGGAACAAA	AGICIGGCTT	CATTCTATGA	CCCCCACGTT	GAGTTTCTTA
199921	ACTAAACAGT	CTTTTCAATT CTAAATATAA	AGGAGIGICC	CONCERN	CCATCAGACA	TAACTAGCCG
199981	CCCATCATTT	AATTTTTCT	ACTCCCTTAT	CTACTCCTGC	TGAAAACATT	TTAATTACTC
200041	CACAAGAAAA	CCTGGCATAT	ACIGGGIIAI ACATCCATTC	AACTTC	AGAGTTGGTC	TTGTGTGCAA
200101	TGTACTATTT	CATCTATTCT	TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	MAGIGIATGC	CACGTGCATG	TATTCCTTCA
200161	AAAAATGAAA	CATGTATTCT	TCACAIC	TGTTTTTTCC	TCTAAAATTT	ATTTCCTTTT
200221	ATTTTTAAAA	ATTTTGCATT	TGACTAAATT TCTTTTCTCT	TGTCAAATTT	AGTCAAATTT	GTTTAAAACC
200281	ATTTTTCATG	TGACCTCAGT	AGIIIIGAGI CCXCTCCTCT	GAAGTTAGTA	CTTCAGAAAA	ACTGTTTTGT
200341	GTTTTGAGGA	TGACCTCAGT	CCACAGCIGI	GCATTTCCAT	TTCTGCGTCC	ACACACATTT
200401	ACCGTCATGA	AATATAGGAA	CGACAAGAIA	AAGTTCAAGC	TCCTGGACAT	TGCATAAAAG
200461	TTTTGGTTTG	CCTGGTCCTG	TIGNCTICCC	AGATTTCCC	GCTATTTCCT	AAGTTGAGAT
200521	ATACAGTAAA	GATGCTTTGT (	GIIIICCIMA. Pycycyczana	MATCAAAATA	GGTTTTTGCC	TTTTATGATT
200581	GAAAGTCAGA	TTCATCTAAA	TIGIGIGAMA	CCLCACARMOS	TACAAAAAA	ACCTAAGGAA
200641	CTCTATCTCT	CTCTCTCAAT	TELECTIONS	TOTA COME	ACTACCTTAG	TTATTATTTT
200701	GTATGTATAT	CTCTCTCAAT (	OIMINITIUG	CTCCLTCTATA	GGGTGTGTG	Tagtgtgtgt
200761	CACTACAGTA	ATCTGTTTCT	TOTAL TOTAL	GIGGATGTGC	ACAACGCATC	CTGCTTTGTA
200821	CACTACAGTA CTTGTATATA	TACACATIII	LYCINAIGINA .	TICAATATTG	TIGAAAACAT	TTTAAAAAAG
	CTTGTATATA		ACACATACA	IGCATGTATG	TACATATACA	CATACAGACA

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200881	AAAATGTATO	CTATGTATAT	TCACACATGT	ATACACACTO	ACACGTACAT	AGAGTTTTAC
200941	ATCCATAGTT	TATAAATGT	GCTTTTTTT	GGTCACCTTT	TTGCTAAGTC	TTACACTTTT
201001	TTTTTTTTT	TTGAGACGG	GTTTTGTTGT	CATTGCCCAG	GCTTAGTCCA	GTAGCGCGAT
201061	CTCACCTCAC	TGCAACCTC	ACCTCCCGGG	TTCAAGCGGT	TOTOTOCO	TAGCCTCCTG
201121	AGTAGCTGGT	ACTACAGGTO	TGCGCCACCA	TGCCTGCCTA	ATTTTTTTTT	TAGCCTCCTG
201181	GAGACGAGGT	TTCACCATG	TGGCCAAGCT	GGTCTGGAAC	TOTORO	AAGTGATCTG
201241	CCTGCCTCAG	ATTCCCAAAG	TGCTGGGATT	' ACAGATGTGA	GCCACTCCACCIC	CCGGCCAAGT
201301	CTTACACATO	TTTTTTTAC	CACTAAACTC	תפתפתום ומפתחה	CCTCATTAGCAC	CAAGTCAACA
201361			מממדבת בידים מממדבת מידים	CDARCOCARA	CREATAACC	ATTTTCTTGA
201421	TTAATATTCA	GAAAAAAATG	GAGTCCCTTT	TATACCACCAC	ACTATATAGA	TTACTCATTT
201481	AAAAATGTAT	TACATTATAT	GAAAGTATTC	AGGTCNANTC	AGIAICIGCI	GATTCATTCT
201541	TTTTAACTGT	GTTATTTTTC	TGCAATGACT	AUGICAAAIG	ARCERCACE	TCTTCCACTG
201601	ATGAAAATTT	GGGCTATTTC	CAGTTTGTCT	TOTAL CACA	MAGIACTCAG	TCTTCCACTG
201661	ACTCAATGTG	TTTACTAATT	TAGGAAGAAT	CARTITIC	TATCCTCCTC	CTTCTCCCAT
201721	TCAAGAATAT	AGCATATGGT	' ATACTATACT	ACACTACTIT	TATGGTATTA	GCCAGATCCT
201781	GTTTTCTGCC	CTTTAATAAA	ביים האתינית היים היים היים היים היים היים היים הי	TTTCTCCCCT	GITTAATTTA	GCCAGATCCT TTTCCTTGTT
201841	CATATAATTC	TTABBBBBBC	TATICIAICAI	ATTOTALOCC	TGAGTCACAT	TTTCCTTGTT
201901	TTTCTATTCC	TGTCTAGTTA	TATAGITITC	MITCHAAGGG	AACATAAAAA	CTTCTTTCCA
201961	ATCTTTCCAG	TCAGTTCACC	, VLICIMCIWI	TOGGAAAAGT	AACTGTTAAA	CCCAGTCATG
202021	TTGAACACTT	CTTATTCCTC	DCACCAAGCC	TCAACCCTTG	TACTTTAATC	CCCAGTCATG
202081	TCCCCTGTAT	TACTGACTTA	TTCATACCA	1 CAACGGG11	TGCTCTTTCT	GGAAGGTGCT
202141	TGGGAAGCCT	TCCCCTGATA	CCCCTACCAC	CCACCACRO	TGGCGCAGCC	CTGTTCTGCC
202201	ACCTGTGCAA	GTTTGTATTG	TTCATCTTTA	TCATCCTTCA	TCATTTGTTC	CTGTCTCTAT
202261	GTGTGGTCTC	ATTCAGTGGA	CTCTCAACTC	TEATCONOCH	ATGTCATGGG	CTGTCTCTAT
202321	ATAAATTAAT	ATTGTCGGAA	CICIOAACIC	TOTOTAGA	ACAGAAAATT	TCAGATCTTA
202381	AAATATAGTA	TGTTGGCTGG	GCCARIGICA	TCAACCCCC	ACAGAAAATT	TATCAAAAAA CTTTGGGAGG
202441	CCGAGGCAGG	AGGATCACAT	CACCTCACAA	TCAAGCCCGT	AGCCTGGCCA	CTTTGGGAGG
202501	ACCTCATCTC	TACTABAAT	ACAAAAAACTA	CCCLCCCCCC	AGCCTGGCCA GTGGTGCCCA	AAATGGTGAA
202561	CAGCTACTCA	GGAGGCTGAA	CCCCCACCAC	CACCAGGCGTG	TGGGAGGCAG	CCTGTAATCC
202621	GAGCTGAGAT	CATGCCACTG	CACTCCACCC	TECECERARE	TGAGACTCCA	AGATTGCAAT
202681	ATAGTAATAA	TAATAATAAT	AATTGTATGG	A TOTTCA A CTC	CTCTGATTGG	ACTCAAAATA
202741	TTTTTAAAA	חידים מידים מידים מידים מידים מידים	TTTTNACTTCC	TOTOTACIG	TACAGGATGT	AAATAGCTGT
202801	TACATAGGTA	AACGTGTGCC	ATGGTGATTT	GCTCCACCTA	TCAACCCATC	GCAGGTTTGT
202861	TAAGTACAGC	ATGCATTAGC	TOTTTTACCT	A A TOTTOTO	CACACCCCCA	ACCTAGGTAT
202921	CCCCAACAGG	CCCCAGTGAG	TOTTTIMECT	CTCCCTCTCTC	CACACCCCCA	CCCCATCCTC
202981	GCTCCCACTC	ATABGTGAGA	ACATCACCTC	TTTTCCTGTGT	TGTTCCTGCC	TCATTGTTCA
203041	ATGTCAGGCC	AGAGAGGCTT	ACAIGAGGIG	CCAMOMORO	ACTITICTIC	TTAGCTGTTA
203101	TTGATGTTTA	TAAATGTTAC	AMMITITIMA	ATTTCAMENA	ACTITICTIC	TACATTACTC
203161	ATTTAACTGA	GTTAACTTTG	TTATATCANA	ATTICATIAA	GGAGTGAGGG	TATTGAGTTG
203221	GCTACAGAGA	TCTTGATTGT	TGGTGGTGAA	GCAATGCAAG	AATTCATTCA	GGTTAAACCA
203281	TAATGTTTAT	TAAGCGTGTA	CTGTCTTAGT	CTCTTCACAC	TGCTGTAACA	TTCAGTAAAC
203341	AACTGGGTGA	CTTATAAACA	ACAAAAAATT	TATTTCAGAC	AGTTCTGGAG	AAATATCATA
203401	TAAGATTAAG	GCCCTGGCAA	ATTTAGTGTC	TEGTERECAC	AGTICIGGAG	GTGGGAAGTC
203461	AGTCCTAACA	TGGCAGAAGG	GTTGAATAAA	CTTCCTTCCC	TTTCTTTTAT	CTTTTTGCTG
203521	ATCCTAGTGA	TGAGGTTTCT	GCCCTCATGG	TATAACTACT	GCCCAAAGAC	AAGGACACTA
203581	AATATTATCA	CTTTGTGGGT	TAGGATTTCA	ACATCACTOR	TGAGAGGATA	CCCTCCTTCT
203641	GATCATAGCA	CACACCATAG	GACAGACACT	GTGCCAACAA	TTGTGGATAT	CAGACATTTG
203701	AAAATGAACA	AGATCCCCTC	AGAGAGCTTC	CDDDDTCCAC	CTATAAAATT	AGTGATTCTC
203761	AACAAATTAT	GCAGTTTGAA	AAATCTACTC	TCATACTCCAG	TTGTGGCATT	AIGUTTTTTA
203821	GGCCACTCTT	TCCTTATTAT	ATTAAATATT	TATE TARE	TGGGGGATCC	ACTOTOROGO
203881	ACTTTTTCTA	CCAGAACTGG	TATCAGCTCA	TGCTCTGCC	TATGCAAATT	ARCARACCT
203941	CATACCTTTT	GGGTAAATTA	AGCCAAGAAA	GTTCTCCT	CTTCTCTTTC	
204001	TCTTTCTCTC	TTTCTCTTTC	TTTCTTCTC	TCTCTTT	TCTTTCTTTC	
204061	TCTTTCTTTC	TTTCTTTCTT	TCTTTCTTTC			TITCITICIT
	<del>-</del>				111C11C1T	ICITICITIC

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204121	TITTTCTTTC	TGACAGGGTC	TTGCTCTATT	GCCTAGGCTG	GAGTGCAGTG	GTGCAATCTC
204181	AGCTCACTGC	AGCCTTGAAC	TCCAGGGCTC	AAGCAATCCT	CCTGAGTAGC	TGGGACTATA
204241	GGCATGTGCC	ACAACATCAA	GCTAATTTTT	GCATTTTTTT	GTGGAGACGG	GATCTCCCTA
204301	TGTTGCTAAG	GCTGGTCTTG	GATTCCTGGG	CTTATGCGAT	TCTCCTGCCT	CAGCCTCCCA
204361	AAGTCCTGGG	ATTACAGGCA	TGAGCCACTG	CCCCTGGCCA	TTATAACTAT	TTTCATTGGC
204421	TTATCAGGCA	CATGATAACT	ATAATAAATC	AATAACCAGA	ארדייייידא ארד דמממיייידידי	DARGARGGA
204481	AGGAATTGTT	TCAACTCTTC	CTGCTACCCC	TCTATCCCTC	AAAAGGGTAG	CCTCDATCTT
204541	GTCCTCCAAA	GATATCCATG	TCCTAATCCC	CAGAACCTGT	AAATATATTA	CCTTATATCA
204601	CAAAAGGGAC	TTTACATGTT	TAATAAGTTA	AGAATTTTGA	GATGGGCAGA	TTTTCCTCNA
204661	TTTTGCAGAT	GGGCCCTAGT	GTAATCACAA	GGGTCCTTAT	AAGAGACAGG	CAGAAGAGTC
204721	AGAATAAGAG	AAAAATACTT	CAAGATGTTA	CACTGCTGGC	TTTAAGGTCG	DECADAGECCE
204781	AAGAGCCAAA	AAATGCAGTG	GTCACTACAA	GCTGAAAAGA	DARACRARTC	CATTOTAL
204841	TAAAGCCTCT	GGAGGGGGCA	CAACCTTGCC	AATACCTTCA	יייייייייייייייייייייייייייייייייייייי	CTCANACCCA
204901	TTTTGGACTT	CTGACCTTTA	GAACTGTAAA	TANACCIOA	TITIGGCICA	GIGAAACCCA
204961	ATCACAGTTG	TGGTAATTTA	CTACAACAGC	AATAAAAAAA	ATTITUTET	3C1CAAGCC
205021	GGAGTTGAGT	AGGATAAGCC	TACTCCACCA	CCTTATATATAG	CONCENTRACE	AGAGATCTGA
205081	AGGATGGCGG	AACTCAATTA	AGGNACTOTO	ARCTCATA	CCCACACACACA	GAGACTCACT
205141	ACTITICATITITE	ATAAGGGTTG	CCTCACACTA	CCAACATCCA	GCCAGAGAGG	GAAGGCTCTC
205201	ביים ביים ביים ביים ביים ביים ביים ביים	ACAAATAGGA	CACACACIA	AGAGGATCCA	ATAGCAACCA	CAGTCTCAAA
205261	GGGAAGACAT	GGATGATATG	DARCACTARC	CACCCCTACA	GCCAAGCAGA	AAATGGATTA
205321	GCCAGGGCAG	TCACAGTTCA	CATTONTO	COTOTOGGG	AGGCAGCTTC	CTGGGAAGTT
205381	TAGCTGACTT	AACTGAACTC	CTCARCACCA	AMONACACO	CCAAATGCAT	ATGGAAAATC
205441	CCAATTAGAA	TATGTATTTC	A TOTAL TOTAL A	AIGAACACCT	CATTTATTGA	GGAGCTACTA
205501	COMMITTAGAA	PCCCCARCCC	ATTIGITCAA	TAACCCCATG	AGTACAGTAA	CACAATCCTT
205561	ACACTACIAA	AGCGGAAGCC	AATTCAAAGA	GGTTCAGTGA	CTTGTCCAAG	CTCAGGGAAA
205621	TCCTCTCTCTT	GTGAATATGG	GTCTGACTCC	ATCACTGATT	TCAGGAGCCC	TGCCCTTTCC
205681	CACTTCAAAA	GCCCCCTTGC	TTTCAGAAAA	AAAGGCTTGT	TGACTGAATG	GTTGTATGCA
205741	CAGIICAAAG	CAGAAACACA	CGATGACATC	TTTTGAGATA	CTCTAACAGT	GAGAACTTGA
205/41	AAATGAAGTT	AAAAATTAAG	CGGCAAAACC	AAGCCGAGGC	TTTCTGAGAA	AGTGGGGCCA
	AACCIGITGC	CGTCTGACTG	CCACGTGGCT	CACTATTTAT	CCCTGTAAAA	ATCTGCAAAA
205861	GIATITGAAA	GGGAAGAAGG	GACAGAAAAC	TCCCTCCTTT	TCCAAGTTAG	CCTTATAGTC
205921	TAGGGCTTAA	AATACTGGTT	TAATGGTGAA	GGTAAGTGCT	TITCTTCTTT	TTGGGTAGAA
205981	GGATTATTAC	TAACTTACCA	AAGGTCCATT	AAGGGGAGGG	AACAGTTTTA	GGAGAAGTCA
206041	GAGAAAAGAC	ATTAACAGCA	ACATAAGGAT	CTCCATCTGG	TAATATTGCC	TAATTCCAAA
206101	ATGAAGAGAC	TCTCTGAAAA	AGATAACTGA	TTCAATGAAG	ACCCTAGGGC	AAGGCTTGAG
206161	AAGCCACTGG	TACCAATGGA	CACTGTGGAC	AATGGTCATT	TCTCCAAGGA	CGCTGTGAGT
206221	ATTAACTGTG	ATGCTGTGAT	TAGTCAGACT	GGGATTGGCT	GTGGAATGAA	ATACTGATCA
206281	GAACTGACAA	GATTTGTGTT	TGGGACTGTG	GCTAACGAGT	CTTTTCAGAC	TTCTATATGA
206341	ATTTGAAATG	GTCTCTCAGG	AAAAGGAGAA	CATGGCCGGG	CCTGGTGGCT	CACGCCTGTA
206401	ATCCCAGCAC	TTTGGCAGGC	TGAGGCGGGC	AGATCACTTG	AGGTCAGGAG	TTTGAGACCA
206461		CATGGTGAAA				
206521	CGGCGCGTGC	ACCTATGCGC	ATGCATAGTG	CGCGTGCCAG	CTATTCAGAA	GGCTGAGGCA
206581	GGAGAATTGC	TTGAACCCAG	GATGTAGAGG	TTGCAGTAGT	TGAGATCATA	CCACTGCACT
206641	CCAGCCTAGG	TGACAGAGTA	AGACTCTGTC	TCAAAAAAAT	ATAATAATA	AAAGAAAAGG
206701	AGAACATGAC	CAAAGTTATG	AATAAGACTG	AAGGCAAGAA	AATTGTACGC	TTGTAGAGAT
206761	CACCTAGCTT	GTTGCCCTCA	TTGTACAGCT	AAGAAAAGGC	ACCCAGGGAC	ATTGTGGTCA
206821	GCACCAATTT	CTCAGAAAGA	TAGGCAGATG	ATGAGAGGGC	CCTCAGTTTT	TCTAACACTG
206881	AAGGAATTGC	TTCTATGTTT	TCTGGTGAAC	TCCTCCCCAC	TCATCTTGAG	GATTCCAGGC
206941		CACTTTAAAA				
207001	CTTTTATAGA	AATACATTTC	ATTTGCTGTT	GGCCTGTATT	TATGGATCTG	AGAGGGCTAG
207061	ACTGCCAATA	TTGTGACTGT	TTATTATTAT	TGCTGTTGCT	AGTATCTAGA	ATATTATACA
207121	ACATATAACA	CTTTGCAATT	TACGAGGCAT	GTCTCATACT	TTTGTTTTCA	CTCCAAACTG
207181	CCCAGTGAAG	TAACATTATC	CCAATTCTTC	CTATGAAACA	GTGAAAGCCC	TAAGAGTTTT
207241	TGAAACTTTA	CCTGGTTTAC	TCAATTTGGG	AATGGCAGAG	CAGAATTCAG	TCCTTGAATA
207301	TCCTCCCACT	GCAGGTTCAT	GCTCTTTGAT	CTAGGTGTAA	CATTTACTCT	GAGTAAACTA

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207361	GGACTCTGGG	CTAACAGAGA	TGAAGCAAGA	CAGGCTGGAT	ATTAGGAGAA	TCTAAGAGCA
207421	ATCTAACGAC	CATTATAATA	AAATCATGAG	TTCTAGACTI	AAAAAAAGGG	AAAAACCTGT
207481	TTTTTTGCTT	ATGCGTATAC	CATAATATTT	ACATTATTTA	TTTTTTTCTC	AAATTCAACC
207541	TATACGGTGT	CAAGTAATTI	TTTTTAATAT	AACATTTTCC	TTTAACTTAA	TTTCAATTCA
207601	TTTTTCTGTG	TCTACTTACA	ACTTTGGCAC	TAGAATTCAC	AATTTTTTTT	TAGAGGTATA
207661	TCTCCTTAAA	GGGAAGGGTT	CTGACACTGT	TACATGTTCT	CAATTGTTTG	CABATAGGTT
207721	AATAATTATT	CCAGTGTCTC	TAAGTACATA	TCAACCATGO	CAGTGTTCAG	CCTCCATAAT
207781	TTTATTAGCT	TCTGTGCTTA	TTTTGGAAAA	ACATTTCCCA	TTACCATGAA	AGACCTCAGT
207841	TTAGGATGGT	TTGGTATGTT	AGCCTGATTT	CTGCATTCGT	CTCATGCAAA	GGAAAATAGG
207901	AAACGAAGAA	CTGAAATTAC	CTATTGATAC	AAAATCAAAG	TAGCATTTGA	AACCATAAAA
207961	CTTAAGTAGG	GCTTTTCATC	CTTTCTCGTT	AGACAGCAAC	AGAGAATGGG	AAGAAAAACT
208021	AAAGTGATGG	GTTTGTGATA	CAATTCCAGT	AACATAAAGA	GCAAGGAGAA	GTAGTTTTCT
208081	TGTGTTTATG	TTTAATATTC	AAAGCTCAAC	CTAAAAGTAT	TTTTCATTAT	CAAACTTCCT
208141	TCTAGAATAA	ATGATTAAAA	CTTGATTTAA	AATATACAAA	TTCTCCTTTA	TARTACCTCCT
208201	AAATGGAGCT	ACCCCATTGA	GTTTTAAGCT	TGTGATTAAA	ATATTACGAA	AACAAACCCA
208261	AAGTTGTAAT	AGGTAGAACA	AGCAGTAGTC	TAGGCATTAG	GGGATCTGGT	CCTCCCTCTC
208321	TGCATCATGT	GGTTTCAGGC	AACTTTTCAA	ATTTTCTACG	CAAATTTTCT	TATCARTARA
208381	ATAAACAGTT	GGGCCAGAGG	ATCTCTGAGT	CTCTTTCAGC	TTTCAGTGTT	TATCAATAAA
208441	GAGAAGTTGG	TGGGAAAGCT	TTAAGTGGAG	TGTAAGTAAT	TGCAGCTGCA	TOTAGACTO
208501	AAGAGTTGCC	TTCAGCCAAG	CCACGGGATC	TTGCATAAAA	AGTGAAATCA	ADTRONANT
208561	GGTCCAAACT	CTGGGTTTGA	CCACAGATGA	CTTCACCTAC	GATCTGAGTG	TACACCAATC
208621	AGCTGAACTC	CTGATATCCA	GATGTTAGCA	AGACTTGGAG	GCCTTCTAAG	CCACACCAAIG
208681	AACCAGTATC	TGTCCTGGTG	CTGACCTGAT	CTTACTAGCA	ATTGGGCCTC	CATTOTOCCOMO
208741	CATTGTACAA	AACAACAACA	ACAACAACAA	TABARTCTCC	AAACACCCAA	AATTGGGTC
208801	TTAGATGGAG	AGATACTATT	CCCAGAATTC	TAGAGATATT	TGGAAAGCAG	AATICAAAAT
208861	TTGCCATGCT	GATGAAGTCC	AATTATTGCT	CTTTTAAATA	CATTTAGCTA	CHICAGA
208921	TAAAATGAGT	ATCTACTAAT	TATTTACAAA	ATCACTTCCT	AAATATAGAA	DOMORGRANA
208981	AATGAAGTGA	TCATCCTGTT	TTCTDACCCA	GARATAGTCA	TTACTGGCAC	AGICACAAAG
209041	CAGTTTCTAT	TCCTGTATGT	GGATGTGCAC	ACCCTATCCT	GCTTTGTACA	CENCERCE
209101	AGCATTTTTC	TAATGTAATT	CAATATTCTC	CANANCATOR	TAAAATAGCT	CTAGAGTACT
209161	TAATCTATCA	AATTGACTTG	CCAGACTCTC	PALE PART CALL	TAATTTATCT	CELLCATUACAA
209221	GCAGTCATGA	GTAATACTAC	DANGGATATT	TTTCCACACA	ATTTTTCATC	CTAACATTAT
209281	TTTATAATCC	TTCATCCTAA	GGTCACAGAT	TATCAATATC	TTTAAAGTAC	TATGCCTTTC
209341	TTTAAATTTT	GTGTGCAAAA	ACACTCCAAA	CCCTTCAATAC	ATAAAATAGA	GGACAAGTCT
209401	ATGTGTTTTT	TTGTTTGTTT	GTTTTGAGAC	CCATTCCTCC	TCTGTCCCCC	GGTTTGATAT
209461	GCAGTGGCAC	GATCTTGGCT	CACTGCAACC	TTTCCCTC	GGGTTCAAGC	AAGCTGTAGT
209521	CCTCAGCCTC	CTTAGTAGCA	GGGTCTACAG	CONTETECT	CCACACCCGG	AATTATCCTG
209581	ATTITTAGTA	GAGATGGGGT	TTCACCATCT	TEGECONGENT	GATCTCGAAC	CIGITITITGI
209641	AAGTGATCCA	CCCACCTCAG	TATCCCAAAG	TGCTGGGATT	ACAGGTGTGA	ACCIGACCIC
209701	CCGGCCGATA	CATGTGTTTT	TABACTCACA	CABATTTCAC	ATGTCTTGAA	CCACTGCAC
209761	CAATTTAAAA	AATAAAGTCA	TAGAAGCTTC	AATTTAGGAA	TGAATGGAAA	GGATTTTAAG
209821	TTCTTAGGAT	ATGGATTTTT	CCTADAGGAA	ACTITAGGAA	GCATCCCCAA	ATTGATGATA
209881	ATTAGTATAC	AAATATTAAA	TTAAACATGT	CCATATTAT	AGCCATGAAT	AGATAATTTG
209941	TGTCACAATA	GCTGGATTTA	TTCACAATTC	TACTANTTAC	TCCCTGTTCA	TELETITICE
210001	CTAGGTGATA	TGAAGACTTT	GTCAGTCCAA	GCA ACTOTOC	ACATTGTGTG	TIMIMATITI
210061	GAGAATAAAC	ATTTTAAACT	TTTAAATCTA	ATACATATTA	GTGTTATGTA	AMOTGAMOGE
210121	TCATGTTCGA	AGGCACATGG	AACATTGTTC	TGGTGGTACA	GAGGGGAGAG	AIGICAICCI
210181	AGAATGAAAG	GAAAGACCGC	TCTGGAACCT	TCCTCCTTAG	CTCTTGAGCT	TACTTER ART
210241	GTCCTGTCTT	ATGGTCTGCT	ACANGCAATA	CCFCCTIVG	CCTTCGCATG	ANGITIAATT
210301	GTTTGATAAA	GTACATGCAA	Thirth C July	DATTOTOTION	GCTGCACTAA	CITCICICIO
210361	TTATCTTTAT	TGAACAGATG	AGGARATGAA	TGATTAGAGA	ATTTAAATGA	CTACCTCTAC
210421	GTCACACAGC	TGGAACTTAC	AGCCAGATTT	CCTTTTAACA	PACCACLFF FOR	CIMUCICIAG
210481	CCAGTAGTGC	CCCATAAAAT	GTAAGTTATA	GAGCTGTGTT	GGGTCNANG	TOTAL COLOR
210541	GCTAAGAGGA	GGCAACATTA	ACANGGGGAN		TO TOTAL CARRANC	TITIMCIGMI
				11910		TOTALIMEDI

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210601	TCTCTCCATA	GATAAAAGAC	TGTCGTAGTA	AAAGAGATTC	AGGGCACAGG	GAAACTCCAC
210661	CACAAAGCGT	GGTACCATTT	CCCACAGAAG	CTAAATGGAC	GGGAAGCCTG	CCACCAGGAA
210721	AGGTAAAGCC	ACTGCTCTTG	TTTGCAGGCT	ATGTTAATAA	GCTGAAGCTT	ATTCCGACAC
210781	ATTTACACAT	CTCTGCATCA	CACTGACCCT	TCGTAAAGAT	ACTCCCAGTG	TAACATTGGA
210841	GCCAGCTCCA	GCCCCTGATC	CTGTTGCTTT	TTCCTTAGCC	CCATGAAATC	ATCTGCGAGA
210901	AATTAAGCCA	AATAAGCAAT	AAATCCTGGG	ATCTAGGGAG	TGGAATAAGT	TTTGGGAAAG
210961	TCTTTTTTT	TTTTTTTTTG	ACTGAGTCTT	GCTCTGTCTC	ACAGGCTGGA	GTGCAGTGGT
211021	GCGATCTCGG	CTCACTGCAA	CCTCTGCCTC	CCGGGTTCAA	GTGATTCTCC	TGCCTCAGCC
211081	TCCCGAGTAG	CTTGGACTAC	AGGCACACAC	CACCATGCCC	AGCTGAATTT	TTGTATTTT
211141	AGTAGAGATG	GAGTTTCGCC	GTGTTAGCCA	GGATGGTCTC	GATCTCCTGA	CCTCGTGATC
211201	CACCGGCCTC	GGCCTCCCAA	AGTGCTGGGA	TTACAGGCAT	GGGCCACCAC	GCCTGGCCCG
211261	GGAAAGTCAT	' TTTAAACCAA	CCTATGTATG	AATCCCTACT	ATAATATTCT	CACCAAGCGG
211321	CTGGCTCTTT	CTCCTGAGCT	TGGAAACCTC	CAGTAAAATG	GAAATAATTA	TTTCCCAGAC
211381	CACCACTCTT	' ATCTGTGAGC	TTTTTTGGCC	ATTAAAAATT	ATTTCTTCCA	TTATATTTTT
211441	ATCTGTGTCT	'TCACAGGTTT	TCTCTTTCTT	TCACTTTAGT	GCTTTTCTTC	AAATAAGCAG
211501	GAAAAATCCA	ATCTATCATG	CACATGGGAA	CCCTTTCAAT	ATTGGTCTGT	GGTTGTTCCA
211561	TTTTATGGGG	ATGCTTTTAA	AGAAAAAATT	TGTCCTTTCA	ATATATTGAA	TATCTTCCAG
211621	CACCACATCA	. CCTGCAAGCT	TTGTAAAAAT	AGTTCTACAT	ATTAATTTTT	D.L.L.L.L.L.L.L.L.L.
211681	AGATTGAGTC	TCATTCTGTC	ACCCAGGCTG	GAGTACAGTG	ACATGATCTT	GGCTCATTGC
211741	AACCTCTGCC	TCCTGGGTTC	AAGTGATTCT	CCTGACTCAG	CCTCCCGAGT	AGCTGGGATT
211801	ACAGGCATGC	ATCACCATGC	CTGGGTAATT	TTTGTATTTT	TAGTAGAGAT	GGGGTTTCAC
211861	CATGTTGACC	AGGCTGGTCT	CAAACTCCTG	ACCTCAAGTG	ATCCACCTGC	CTTAGCCTCC
211921	CAAAATGCTG	GGACTACAGG	CGTGAGCCAC	TGCACCCCAC	GTAGTTTTTT	TTTTTTTTTA
211981	AGTTGAACAT	ATGTGAAGGC	AGGACCTAGT	GACACATAGC	AATAACATTT	CCAAGTAGAC
212041	ATTACACTAG	GGAATTAGTC	AAAGTGCTCA	TTTAAAGTAC	CATCTCTCAA	ATGTATTAAA
212101	AGAGAATCCT	TGGATGTGCA	ATACCTTAAT	TCAAAGGCAG	CTCGTTATGT	ATAAACTCTC
212161	AAGCTTTGTG	ATAAACAAAT	GTGCATAACA	GATGGGACTA	TTGACTTACA	GCCCAGGGAA
212221	TTTTATTGAC	GCTGAGAAGG	TTATGTGACT	GGCTCTGCCA	CTGTCATCCC	CATTCACTTC
212281	ATTTTGGAGC	AATATGACAT	AAATGCCTTA	CATGTGGGTT	TTCTCTATTT	ATCATGTGTT
212341	TCCTATCCCC	TTGAAAGATG	GCCATATTTG	CTTTACTTGG	TTATAAGATC	CCATATTCGC
212401	TGTCTTGAAG	CCAACCAAAT	AATTTGACAA	AGTGGGTTTG	TAGTGCTGGC	TATTTTGGTG
212461	AAAAAAAGAC	AATGAGACTT	CATGTGTCAT	CCAAAGTTCT	ATCAGATCGA	GCTGTGAGAG
212521	AAAGGAAAAG	AAAGGGGTCT	CAGTCAGGAT	GCTCACTGCA	TACATCTGTG	TTGTTGTCTA
212581	GGTCCAGATT	TCTGTTCATT	ACGCTATGGG	CTGGCTCTTA	TCATGCACTT	CTCAAACTTC
212641 212701	ACCATGATAA	CGCAGCGTGT	GAGTCTGAGC	ATTGCGATCA	TCGCCATGGT	GAACACCACT
212761	TCCACCAMA	GTCTATCTAA	TGCCTCCACT	GAGGGGCCTG	TTGCAGATGC	CTTCAATAAC
212821	TOTTON	CCATCAAGGA	ATTTGATACA	AAGGTAAGTA	TGATGGAAAA	TAGGGCTCTT
212881	ATACATTECC	AAAAACTTTG	AAAGGAAGGC	ATAGATCTTG	ATTCTGTGGA	GTATGGAAGT
212941	CTTCNATANT	AATGACAAAT	TAAAACTGAC	TGGAACTATT	TTTCTTTGAG	ACATTGCTTA
213001	ACACTTANA	AAAAATAAGA	TTTCATTGAG	GTTATTATGA	TTATAAGGTG	GGGGAACTGT
213061	TAGGTATTAG	GTGAAAAATT	TAAAAATGGA	ACAGTTTATG	TGATGTCTTC	AATGAAAAAC
213121	TTATTCTTTC	CTGGGCACAT	TCTTATAGGT	TACTCAATCC	TATTCAGTTC	TCTGCCTGTT
213181	CGATTCTTGT	TGAGCAATTT	CCARAGERA	TAAATTCTAT	ATAACCAATA	GAAATGCAAA
213241	TCTCCACTCA	CCATAGCTTT	CARRIAGE	TTTGCCAAGA	GAAAAATCAG	TTAAAACTTT
213301	TGAGATAGAG	CCTCCCAGTT TCTTCCTCTG	TCATTAGCCA	ATTTTGCTGT	TTGTTTGTTT	GTTTGTTTTT
213361	GCAGCCTCCG	CCTCCCGGGT	TCATICAGGC	TEGAGTECAG	TGGCATGATC	TCAGCTCACT
213421	GTAAGGGGGC	ATGCCACCGC	CCCTCCCTA A	TICCIGICIC .	AGCUTCCCAA	GTAGCTGGGA
213481	ACTAGGCTGG	TCTCGAACTC	CTCLCCTCLC	CTCATCCA CC	TITAGTAGAG	ACAGGGTTTC
213541	TTGGGATTAC	AGGTGTGAGC	CACTGTGCCAG	GIGATECACC	TATATOTA A A	COCCAAAGTG
213601	GCATTGCTTC	CTGCTTGTGT	TATGCGTGAT	TCTTCIGCIG	₼₼₼₼₼₼₼₼ ₮₩₩₩	GICIMITICA
213661	CATCTTACTT	ACTTCCTCCA	TTAATCAATG	AGTTAAATAA	Y Dutchturians	CCAGIIAIAA
213721	TTTACATTTA	TATGAAAACC	ATGAATTTAC	CCAATTAAAA		GIWIGIIIWI
213781	TTGTACTGTA	CATTTCCCAT	GTCATCCCTA	TAATTCATGA	ጉጉልልጥርልምም ፡	110001101C
		<b>-</b>				TUTUTUTIO

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213841	GACCTAGCTT	ATTTACAATG	AGTACATAAA	TTTATTGTCT	CCAGTCTTTC	CTCCATTATC
213901						CCTTCAAGTT
213961						CCTGCTTCTC
214021						CAAGCCAGAG
214081		TTTCTTCCAA				
214141	AAATTTTCCT	TTTTGACTTT	CTACAAATCA	TAGTCATTTT	TCAAGGGATA	GTTCAAGTAT
214201		CTGGGACCTT				
214261		CCTCAAATCT				
214321		AGGTAAAATA				
214381		ATTTAGGGGC				
214441		TTATTTATTT				
214501		TTATTTAGGA				
214561		AGAAGAATGT				
214621	CAGCCAGGGC	AGACCATAAT	TOCONNACTO	CY CALGRAM CAME	TOTALITIES	GAACICAGII
214681		TCAGTAAAGT				
214741		TCAACTTTTA				
214801		TTTTCTTCCA				
214861						
214921		CTTCATTTCT				
214981		TTTGGGAAGG				
215041		ACTAACCTCA				
	CTTGACAAAG	AGTTTTCCTA	TTTTTCCCCC	AGGCCTCTGT	GTATCAATGG	AGCCCAGAAA
215101		CATCTTTAGC				
215161		AGGGATATTT				
215221	CCCTTCTCAC	CCTCTTTACA	CCACTGGCTG	CTGACTTCGG	AGTGATTTTG	GTCATCATGG
215281		CCAGGGCATG				
215341		ATTCTACAAA				
215401		AGTTTACTAT				
215461		CAGGATCAGG				
215521		GTCTTATCTT				
215581	TGCTGAAAAA	TTCAACAATA	TAAGACACTT	GCATCACAAA	TAGGAAAGAT	GCATCTGTGC
215641	AGTAAAGACA	TTGAAGCTTA	GAAGTAGAAA	AAACCATTGT	GAGCTAGGTT	TCAGCTCAGA
215701		TAGTCAGAAA				
215761		ATTGCACACA				
215821		AAGTGAGAAT				
215881		TTTGAAGTTT				
215941		ACAACAACCC				
216001	AGGGAGTTAA	GGAAGGTGGA	GATTAAGAAA	ATTGCCCAAA	TACAAATAGC	CAGCAGGTGG
216061		ATTTAAGCCC				
216121	TGCTAGACTG	CCTTTCCATG	GTATGTGATC	CTACTCAGGC	CTCTACAGCT	TTATCATTGC
216181	TGTTCTCCCC	AGCCTGTCGT	GCTGAGAGTA	TATACTCGAA	GAGCAGAACT	AAAATTCCAT
216241	CCAGCTTCTC	ACTCCTAGGT	CCACTACACA	GCTGCATCCT	GCAGACTTTT	ACCTCAAGCA
216301	ACCCTCCTGC	GTTCTTGCTT	CCTTCCATCA	TAGTTGTAAC	CATCTCCTCT	ATTTGCAAAT
216361	ACTATCTGCT	GATCTCTCTC	TTCTAGACTG	GTTTCTTTCA	ACCTTCTTCC	CACCAAAACC
216421	AAGTTAGCTT	GCTAAAATAA	AGATGGCGCA	TTTTTACTCA	CCCGCTTGAG	AATTTTCAAT
216481	GTGTTCCTTC	ATGCTTACAG	AGTAAAGCCT	GACCTCTTTA	TTGCATGAAT	ACAAAAGTTC
216541	TTAGCCATCT	GGCCCCAACC	TTGTTCCACT	CAACTCCCCT	GTGCAAGCAT	GGCTCCAGTG
216601	GCACTGGACA	TTGGCTGCTC	TCCACATAGA	TCTGCACTGC	ACTTCCCTCT	GGCTCTGCTC
216661	CCGTTAGTTT	ATATGCCTGG	AAAGTTCTTT	GCCCCTGTTC	CTTGTGCCAA	AATTCCATCT
216721		ATAGCTTATG				
216781		TTTTTTGAGA				
216841		TCACTGCAAG				
216901	CCCGAGTAGC	TGGGACTACA	GGCGCCTGCC	ACCATGACCG	GCTAATTTTT	TGTATTTTA
216961	GTAGAGACGG	GGTTTCAAGC	CAGGATGGTC	TCAATCTCCT	GACCTCGTGA	TCCGCCCGCC
217021	TCGGCCTCCC	AAAGTGCTGG	GATTACAGGC	GTGAGCCACC	GTGCCCGGCC	AAAACTTCCT

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217081	AAATCTTATA	ATTATTATCA	ATTTATCCTC	AGATATACTI	CCACGTACAT	TGTAGTTTTA
217141	TTATATTTAT	ATTTTACATO	TTTTTTTC	AATTGCAGTT	TGGGACCCAT	TAGTGAGTCA
217201	TAAAATCCAT	'TGAGCGGGTI	. AAAATCATTA	TTTTAAAAAA	TGAGTAGAAT	' AGAATAGAA
217261	TTGTTGGAGT	GCATTGGACA	TGGTAAAGTT	AAATATCGAT	TCATGAAACC	ATCGTTTGAG
217321	GCATATGTGT	GTGGTTGTAT	GTACAAGTGT	TTATGCATAT	TGGTGTGTGT	GTTATGTTAC
217381	CCTGTAAAAT	GCATTTCTTA	CTATAGGTCT	CTGTGAAATA	TGTGTCTTGT	TGTTTTTTAA
217441	TGTAGACTTC	CAAAGCCTAC	ATGGCATTTC	ACTAGTGACA	איידידים במרדים ב איידידים במרדים	TTCACATTTT
217501	TCTCTCCAAT	TGGACCAGAA	GCTCTTTGAG	GGCAGGGGCT	GTATCTTACC	GATTTTTGTA
217561	AGTCTTTCAT	TTCCTGCCCC	TAGCCTCATA	TTAGATCATG	CAAGAATGCA	ACTGTAATCA
217621	CAAGAAAATG	CTAATGGGCT	GTGATAGCAG	AGAGTTACTG	TCACAAACTA	AGGGATTTAG
217681	ATTTGGTCAC	ATTGGTGTTG	AGGAGCCATT	GAAGAATCAG	AGAGTGTGTT	ACTATTATTT
217741	GTTAATTTTA	ATTATATCAT	ATTACTTTAC	TGGGGAAAAT	CTGTGAGCTA	TTTTAGAAAT
217801	AAATACTCTC	ATTGCCCAAT	AATTCTAAGT	CTGCCACCTC	ACTETTECEN	CATTGTTTAG
217861	GGAGGCCACG	AAGTCTCAGC	CTTTGATATT	TTCATAAGTG	TOTALIGGGA	TTTTTCCTTT
217921	AGGGTCAGCA	TTTGGATCCT	TCATCATCCT	CTGTGTGGGG	GGBCTBBTCT	CACAGGCCTT
217981	GAGCTGGCCT	TTTATCTTCT	ACATCTTTGG	TGAGTCACTT	TOTOTOTANA	CCTAATGCCT
218041	CCATTTCCTG	AGCATCCATT	TTGGCACCTA	CACCACCCAC	TOTOTIMANI	TATGAAAGAA
218101	AATGTCCTTT	ATCAAATGGA	AGATGATAAA	AAATGTCAAC	GGTTGGTATC	ATTTTTT ATC
218161	TAGTCACACA	ACCTGATTAA	CACCTTCCTG	GTGGTTCTCC	Chaccoacac	ATTITIAATC
218221	GAGGAGTTGA	CTATTCACAT	GGCACCCACC	GACTTGTGAT	CCNCCCACAC	GCAAAAGGTA
218281	CAAGCACCTT	CTGCAGAATC	TCTACCACCA	CATCTGAACT	CCCTCCTATA	CCTTCCATAT
218341	ATGTCAAAGA	TAGTGAAGTA	CATTTTCAAT	GTGTCTTCAT	ATTTCATTA	IGCAGTTAAG
218401	TCTGTCCAAG	ATGCCTTTCA	CCTGTTCTCT	DCCD BCTTA B	ATTICATIAT	AATTATTATT
218461	ATGTTCCCTT	CCCCATGGGC	CCTTCCAGGG	CTTACCCTCT	CACAMMOMO	TTCAATTCAA
218521	TTTATGATAT	TTCCTCTCTA	GGTTATGTTG	GTGTGTAATT	ATTTATTTCT	CATTCTCTCC
218581	CCACTAGACT	GTGAAATGCT	TGAGGCAAGG	AATCCATTCT	ATTIMITED	COLLITICITY
218641	TCATCATGGT	GCCTGATTTT	TAGCTTTAAA	ATABARCART	CACTCAATCC	CACTIGGGIG
218701	GGGGATTTAA	AGAAAACTAG	TCCTCAGAAT	ענישטענישטעני	AGNATOTO	AGTAATTAGA
218761	AATTCCAATA	ATAAGACAAT	TTTCTACACT	TCATTTTCTT	TOTALIGITE	1 CAAATAAGG
218821	TTAAATATAG	TCCTGGCCTG	AATGGCTTTC	TCATTARTCA	TOTTALAGUCA	AATGGTGTCA
218881	ACATGTTAAC	CAGGTATTGT	ACADADATAT	TTCTTTTCCC	AATCCATAAT	TITGGTTTGT
218941	CTTGAATACA	AATAATACTG	TCTCTTCTA	GTGCTTTTGG	AAICCAIAAI	GGATGTATGG
219001	TTTCATGGAA	GGTTGTTTCG	TGTATGTATG	ACTCCARACC	TCR CMR TWO	TGCCACATGA
219061	AACAAGACAA	CTTATGTGTG	CATTANGANG	TTGCTGCCTA	AAACTATTCA	GATCTTCCGC
219121	ATTGGAGACT	TTAAAGTAAT	TAATCAGCTA	TECANTECEN	CGCTCCTCTT	CACTGTAATC
219181	GGCTCTGACA	TTGACAAATG	GTGGCTTTCT	ATTTGACACG	TARTATOTA	ATCTCCAGAG
219241	CAGGTTTGTA	GAAGGATTGA	AAGAAAGAAT	GGGDDCDTTT	DCCTCCTTAN	AAAGCTTTAA
219301	GCATTAATTG	ATTAGTGTGT	AGAAGGGAGA	GGCATGCCAC	TTCACACCAA	GGTAGAATAA
219361	CCCAGTAAAC	AAATCTACCT	AAAAACTAAT	TTTATCCCTT	CTTCCCACCT	ACTICCTICC
219421	GTGTCTGCTG	TCTCCTATGG	TTCACAGTGA	TTTATCATCA	CCCCAGGI	AGCACTGGCT
219481	TAAGTGTTAG	GGAAAAGGAG	CACATCCTGT	CCTCACTGGC	TCAACACCTA	CACCCGIGCA
219541	CTTGTACCTG	TGGCCCATGC	AGAGGTCTCT	AGGGCAGGGT	GTGGATCTCC	TOTOLOGO
219601	ACCATCTTGG	CTGCTCTAAT	ACTCATGCTG	ATTAGATOTT	TOTAL	CCLCMMOMOGC
219661	TGGACGAGCT	GTCCCCATAA	AGGCGATGGT	CACATGCCTA	CCACTTTCAGC	CCAGTTCTCC
219721	GGGTTTTTTC	AGCCATTTCT	GGTTATGCAC	CATCATCCTA	ACATACOTAC	CCATTTTCCT
219781	CAGTACTCTG	CTCCATGTTA	ACATCAGAGA	TOTONOCTON	CONTROCTAC	CAACGTATAT
219841	ATGATAATGG	TAATAAGGAG	AAACAGTTCT	CTCTTACCTA	TTT CATTO	TTCTACGAAA
219901	TAACCATTAA	TTTAACCTTC	ACAATGACCT	TGAGAGAGGC	TACATICIO DALCALICIO	TTCCCTTTTCATA
219961	ACAGATGTGG	AAACAGGACA	CTTAGAGGTG	AGATAACTTC	CCCCFCGGGGG	CACAAMACOO
220021	AGTGATAGAG	CTGCTGCAGC	ATCCATATTC	TTDACCACTA	TCCTATGIIG	CCACACTACTA
220081	TGATTCCAAA	GCTTCTTTTA	GAAATAATAT	TGCTGGGCCCA	GGCATGGTCC	CTCATACCAGC
220141	TAATTCCAGC	ACTTTGGGAG	GCCGAGGCAG	GCAGATCATG	DCCTCDCCD T	TOTALGUUIG
220201	GCCTGACCAA	TATGGTTTAC	TAAATATCAT	CTACTAAAA	TACAGARA	A CCCA CCMCM
220261	GGTGGCAGGC	ACCTGTAATC	CCAGCTATTC	AGGAGGCTGA	GACAGGAGAA	MCCCMGGIGI MCCCMGGIGI
				MULJUULTUM	AADADDAJA	TCGCTTGAAC

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220321	CCAGGAGGT	G GAGGTTGCA	T TGAGCCAAGA	A TCATGCCAC	r GCACTCCAG	CTGGGCGACA
220381	GAGTAAGAC'	r ccgtttcaa	A AACAAAAAA	CCAAGAAAT	ר אמדמדדכרייי	TTATCTGGAG
220441	CCCAGAGTG	A TGCAGCTTC	T GGCCCTCTTA	TCTGAGACAC		GTGTGAAAAA
220501	GGATGCTAA'	TTTCCCCCA	A ACAACCCAC	GTATCATGG	GGTABGTTA	TGGCTGGTCT
220561	GTGTAACTG	A CAAATTTTG	G TGCTAACGT	TCTCTATAA	TACTCTCTA	AAACTTCCTT
220621	CCTTCAGAG'	GGAGTTCTG	CCTCCCTGC	TTTTATTGC	COTGCAAGC	GTACAATTTT
220681	AGGAGGTCA	G CTGGCAGAT	r tecttttgt	CAGGAATCT	CTCAGATTG	TCACTCTCCC
220741	AAAGCTCTT.	TCATCTCTT(	<b>GTAAGGATAA</b>	A GCGTGTGGG	ב ככשיייים אכיכ	2 2 Transcontinuo
220801	CTGCACATG	TCTCAGAGG	TTCCCTGACA	GCATGTCCT	ATTGCCCAGG	GCTCCTCCTT
220861	CCATCAATA;	r GTGCTGTGG(	CCTGCCCTTI	GTGGCCTCC	GTTACCTCAT	מיים מיים מיים מיים מיים
220921	TIGCIGATA	: TTATTCCTG(	GACCAGTAAC	CTATGTGACT	CAGGGTTTAT	CATCAACACC
220981	TTAGATATCO	CCCCAGGTZ	AGAGCTCTAC	CTGTTTTTT	CCCTCCTCC	GACCCCTCCA
221041	GAGGIGITAC	ACCTCAGTG	F TCGCCGTGAA	ACTCTTTAA1	GTTACTGACA	<b>ምምር/ርን ር</b> ምን አመ
221101	GGCAGAATGA	CAAATAACTA	CAAATATCTG	TCTGTGGCCA	עמשליים ביים ביים	AACAAATGTG
221161	GCATTTTTAC	AACAACAATT	TCCAATCTTG	GCCAGTAATC	ATTTTCACAA	AAACCTTCCC
221221	AAGCTTCCCT	' AACAGAGATT	GAACTGTGTA	TGCTGGGAAA	AGGCCCACAC	ACAGGTGATT
221281	I GGAAAAGT I	TCCATGGTGT	TGTTCATATT	' AGCTACCACA	TATATATATA	ጥ አጥ አጥ አጥ አጥ አ
221341	TATATATATA	TATATATATA	ATATATATATA	TACAGTCACA	ATABGCCAGC	TOCTOTOCO
221401	AGACTTGCCA	TATATCAACA	CATCTAATCC	TCACAGTTAT	ATTAGGTAGG	CCCTATTGTT
221461	ATCCCCATTI	TATAAGGGAG	AAGGCTGAGG	CACAAGGAGG	TTAAATGGTG	TGACTATGGT
221521	CACATAAAGG	CAGAGCCAGG	ATTTGGACTG	GGGGAGTCTG	GCTTTGGAGT	CTGTGTCCTG
221581	CCCGTTGCAC	AAACTGGCTT	' CTACACTGAG	CAGCCAGGGT	AAAGAAACCT	COTTCCCACA
221641	GAGACTGCAT	TGCTCCCTGG	TTATTGACTT	GGTAGATTGG	TAATTTCAGG	TTTCCCAAAT
221701	AGACATTGCC	CTGAATGTCT	' TTAGGTGAAT	GAAAAACTGC	ATTAAGCAAA	ATCACTTTCC
221761	CATTAGAGCT	GAATTGCATT	' Aaagttgagt	TGCTGCAGAA	GCTGTAGGTG	CCTTTCTATA
221821	TAAAATCATT	TATAAAATCA	TCTTCCCATA	GATATGCAAG	<b>TTTCCTC3TC</b>	CCAATCTCAA
221881	GGGGATTTGG	GCTCATCGCA	GGAATCATCT	CTTCCACTGC	CACTGGATTC	CTCNTCNCTC
221941	MGGT TGGGTC	AGTTTATTGA	ACATCTTCAA	GTGGCAGGTA	TTGTTTTAGG	TGTTGGAGAT
222001	ACACACGGTG	CTCTAAAGAT	CTGGATGGCA	ACACAATTAC	TCTATTTACA	TCACCCTCTA
222061	AATCAGACTC	TGGTAGGTCA	GATTTCCCAG	AGGAAGAAAA	ATATAAGCTT	ע ערטייטיידאיידער ע
222121	GATGAATAGA	TGTTAGATTG	ATTAAAATGA	GCTGTTCCGG	TGCAGAAGAC	AGCACGTATC
222181	ACTICCIAGA	GGTACATGAG	CATGAAACAG	TTCTTAGTTA	TGACCAGAAT	GAAAGACACA
222241	IGICAAGGAA	TAGCAAGAGA	CGAAGACAGA	GGGGCAAAAG	AAGATCATCA	ስር ስ ስጥስጥ <b>ርጥ</b> ጥ
222301	CAGACTAATC	CAATTTTTAA	AAAATCACAA	AAGGGAAACA	AAGTGTCCTA	CCCCACTTA
222361	AAGATAATTT	AATGTCTGGA	AACAGATCGG	CTGTGAGACA	TTGCAAGGAG	GCTTGCTCGG
222421	IGITTGGAAA	TGCAGGCTCA	TGAGGAAGAT	GAAAAGACAG	ACCCAGGCAG	CCATCCAACC
222481	ACTGACTAGA	ACCAACTTAC	AAAGAGAAGT	TTTGTTTTTA	CTACATTTCT	ATCTCATCAA
222541	GIICCCAGGY	TAATATTTGA	CTAAACTGCT	AGGAATCCAC	TGTGACTATA	ATCCTCCA NA
222601	IGACTTAGTA	GGGCTTTCTG	AGGAGGGTCA	CACAGAAGAC	CARAGAGAAC	TCDTCTTCDD
222661	1 IGAGATGGG	TTATAGTGAT	AGTTGTCAAC	AGCCAATACA	GAAACAAAAA	****
222721 222781	AAACAGCAAC	AACAACAACA	ACAAAAAAA	AAAACAGAGA	ACACACARAC	A C B A TO COOR C
	AAIGCCATTT	TAGGCATAAT	TTTAAATGAG	TAATATTATA	TGTTGAAATC	CAAATTTTTTT
222841 222901	GAMMACATI	AGTGTATTTT	ATTTTTGTTT	AAAGAAATAA	CCATCTCAAC	TCDCDDCCCC
222901 222961	MIGIGCKIII	TGGCCATTTT	GTTTCCAATA	GTTTCATAAA	ت لا لاشتباشياشيا	TA A CITIA CITICO
223021	ACATIGITEE	TTATATTCCT	TGTGATCAAC	ATTGCAATAC	ACAACTGGGA	CCCCTT CTT C
223021	WCIGGIGIA	GAAGGAACIT	GTGAGATTGA	TCATTTTCTC	∪ لا بلادالمالمليليلي	7 TOTA CO 2 TOTA
223061	TIGAGICIGG	TIGGAGGAAT	GTCTTTTTCC	TGTCTGCTGC	AGTCAACATG	TTTGGCCTGG
223201	TOTITIACCT	CACGTTTGGA	CAAGCAGAAC	TTCAAGACTG	GGCCAAAGAG	A CC A CCCCCTA
223261	CCCGCCTCTG	AGGACATAAA	GTTACAAACT	TAAATGTGGT	ACTGAGGATG	A A CHICAMINA A
223261	ACAITITIA	CITCICTCCA	TATTCCTGAC	CATAGACTCA	GCTGTTTTT A	3 CTCTCCCCTC
223321	IGIGITAGIC	TTCCCTGGGG	AGCCTTTATA	AGACACTGAT	ACTTGGGACC	CACTCCACAC
223441	ATTCIGAMIC	AATTGGTCTG	GGGTGGAACC	CAGATACTAC	בי ע הההוהוה עו עוד	3 m3 CMCCMM3
223501	GYGGIIICIM	GCATGCGCCC	GGGGTTGACA	ACAGCTGGAC	AAACTTCAAA	ACTOR BOTTOR
	TGTGGCCTTT	GAAT TTTCCT	CATTGGAAAG	TACTAAATAA	ATAAAAATTC .	ATGTGAAAAT

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223561	GATCACTGAT	AAATATCTTC	ATGGTGGGGC	AGGTTATTGG	ATGCAGAGAA	GATCTGCTCG
223621	GAATTGTAGC	CATATGTTAC	AGATCTCAGC	ACCGATCAGA	ACTGTAAAGC	TATAATCCCC
223681	AGAATTAAAG	TTTTTTATTAT	TTTTTATACA	TTGTAAAACA	TAGACGTTTA	TTTATGTGAT
223741	TAAATTCTAT	TAAAATTTAC	ATGCTAAAAT	AAAATAGACC	ATTTTCAAAT	TATTTAGATC
223801	CAGATATTTC	CATCAGATTA	AACAGATATT	TATTTATCCT	AGCCCAATTG	CAAGAGATTA
223861	ATGATGAGAA	AATGACCAAT	ACAAGATTAA	ATAAATGAGG	TTAACTTAGA	AATCAAGGAC
223921	AGAGAAGATA	GAACTGGAAA	GCTTGTATTG	TGAGAAGAAT	GAATGTGAAG	GAAGGCAATG
223981	TAGACACTTC	CAGAAGGGAT	AGCAATATAG	TTTAGACCAT	ATAATGAAAA	TTGGAGAGAG
224041	ATGACAGAGA	CACTTTCAAG	TGAAATGACA	ATTTATATGG	GGGAGAAAA	TATTGAAGAC
224101	ATAACAAGAT	GAGAAAAGGC	ATAGAAATGT	ATCACATACA	AGGCATAGAA	GTGTATCACA
224161	TACAAGAGAA	GTTCCTTTTG	AGCGTAGAAA	AAGATAATTT	AACCTTCTTC	ATATTTTTCT
224221	TACTTTCCCA	AGATACTCAG	ATAGGCAGCG	TCAACTCTAA	CAGGAATTAA	TTTGGCTCCT
224281	AACACTTAAG	ACATATCCTT	TAGTTTGTCT	CCTCACACAG	AACTGATTCT	GGTTTTGCCA
224341	CAACATGTCT	AGAGAAGAAG	TTCCCACCAT	ATTTTAAATC	ממממדדמד	ACTGCTTGGA
224401	CAAGAACCTT	GGGCTAATTC	AGCAGATGAA	GAGAATCTCC	TAATGCAAAT	CAATGGGTAT
224461	TTTTGAGCAA	GTTTTTCAGA	AAAACAGAGT	GTCAGGCCCT	GAGGGTGGTA	CTAAGATGAG
224521	AACATTGATT	TTGCCTTCAT	GATATTGACA	ACACAAAGAG	GAAAGGGGGT	TTGCAGAAAA
224581	CTAAAAGAAG	AAGTAGAAGA	AAAAAGAAAG	ACATACTATA	ATACCTACTC	AAATTATGTA
224641	CAGAAAAAAG	AGGAAAAAA	ACCDADADAG	GGTGGGGGAC	VINGGINGIC	AAATTATGTA
224701	GGGCCAATGA	CTTGAACAGG	GACTTCATAA	ANGROSSIAN	CTARCTCCC	CCTTAACATA
224761	TAAAAAGATG	TTCAACTTCA	TTAGTCATTA	CAGAAATGAA	AATCAAAACT	CCTTAACATA
224821	ACCACTATAA	AATTAACTAA	TGGATAAAAT	CADAGGAGAT	CCARACT	ACAATGAAAT
224881	ACATGTGGAG	CAACTGGAAC	TTTCATACGT	TACCAATCTC	A A CTTTTCCA A	ATGTTGCCAG
224941	CAATATCTCC	TAAAGCTAAA	TGTACAATTC	CAGTGACTCA	CACATTOTA	AGCTGCTCGG
225001	ACATATACAT	CCATAAAACA	TGTACAACIA	TOTTONTACO	ACCACHARON	TTAGAAATGC
225061	GAACAGGAAG	TTGTCTGTTA	ADDADAGAA	CACTARATAG	AGCACTATCT	GTAATAGCCT
225121	CAATGAGAAT	TAACAGACCC	CARTATATATA	TACAMCA AMC	ACCACGGTCT	ATTTGTATAG
225181	GATTAAAGGA	AGACAAAACG	CAMIMIMIAM	TAGATGAATG	GGTCTCATAA	GCACAATATT
225241	GCTACAACCA	ATCCGTCCTG	TTARRATICITI	TAMAGGITTA	TAAAATACTT	TTTAAAAACA
225301	GTTGTGGCTG	GATGGATGGT	TIMMAMICA	TCCTCCTCCC	CTACTACTAC	AGGGATGGGG
225361	CTTGACTTGG	ATGTGTGTTT	ACTIMISMS	TATTCTACAT	TTATCATORA	TATTTTATTT
225421	TGAATGTAGA	AAATAAAACA	GAAAGCAAAT	TCDBBCTATC	ATCOTTO	CACCTTTA
225481	TCTGACTTCG	TTTTGACCAA	TGGAGCAGTT	GGGAAGGGGT	CTTCCTTTGA	GAGCTTCTGC
225541	CTTTTTTTT	TTTTTTTTT	TTTTAGACAG	AGTCTCACTC	TCTCCCCCCC	CGGICCITIG
225601	AGTGGCTCGA	TCTTAGCTCA	CTGDAAGCTT	TECCTCCCC	CTTCATCCCA	GCTGGAGTGC
225661	TCAGCCTCCC	CAGTAGCTGG	CACTACACCC	ACCTCCCGG	ATTCCCCCC	Trerecigee
225721	ATTTTTTAGT	AGAGACGGG	TTTCACCATC	TTACCCACC	#IGCCCGGCT	AATTTTTTGT
225781	CGTGATCCGC	CCACCTGAGC	CTCCCADAGT	CCTCCCATTA	CACCTCTCAC	CTCCTGACCT
225841	CGGCCCCTGG	TCCTCTGCTT	TCATGTTCTT	CTTGGTCCTC	TTCCTCCTCC	CCACCGCGCC
225901	GAACTTCCAG	TATCAGAGCA	GGAAGGAAGG	Chargeeren	ATCCATCCTCC	TCTTTTGTTG
225961	GATCAAACTG	CAAGTTCTCA	AACAGCAAAA	TTARTCACCT	CICGAIGCIG	1 CAGCTTTTG
226021	ACCCTGAAAG	CATCAGTTGC	TTCCAATTGC	ATCAGTTGCC	ACCCCTCATA	AGAAACCAIG
226081	TGACTCAGAA	TGCCTAGGTT	TTCCCAGCAG	CTTCTCTCAC	COGGIGAIA	CACCETTOTICE
226141	GATTGATTCC	TGACAGATGA	CTTCGGTGTG	TCACACTTTC	ACCOMPANCEME.	CAGCIICICI
226201	ATGGTTTGAG	GAAGAGTTAC	CATTCACATT	CCTAATGGCT	TCAGAATAGA	TCCTIATGIG
226261	AACTGATAGG	AAACATTTCT	AATTCATCTC	CCCTCCCCAT	CCCTTTTCCT	TGCAATIGIG
226321	CAATAGTCAT	GAAAATTAAT	TCVCALANGLA	COLICCOMI	TTTCTCTTCT	ACCENARCAE
226381	GAGATGACTT	ACTTTTTCTC	CTTGACTGTT	TITE TARABLE A	WITCHTCH TOTAL	ACCIAAIGAI
226441	TAATGTTGAG	CTTTCCCTTG	AATATTCTTT	TCATCTACCA	CDCDDALLACS CHITATATIA	TTCT CTAITICT
226501	GTTTATTTAG	GACTTTGGCT	GATGTACTGA	TATATCACAT	TCCCTCTCTT	TICACIANIA
226561	TGTTTTGTGT .	ATCTTTTTTC	TGTCTGCATA	TGGAGCTTAT	COTCATOTA	AAAACAACATG
226621	AGGAGAACTT	TCCTTTTTCC	CCATTACTCT	CARALLIMI	GCIGATITUA CDCTDCDDTC	AAAACAAGAA Caaammamam
226681	AATTGCTGTT	GTTATTTGAA	AGCTTGDADG	COMMUNICATION I	TRARRATG	GAALLITTAT
226741	AGCCATTTTG	AGGAGACTTT	GATARCTOTO	<b>₩₩₩₩₩₩₩₩₩₩₩</b>	TORGIONA CITO	CTCTTTTT 3 2 C
					TOMOTIMOTO	GICILITANG

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226801	GGGTTTTATA	TTTTTTTTT	ATCAATTTT	ACCATTTATO	TTATCTTGGA	GGATCATCTA
226861	TTTTACACAC	TATTTAAAGT	TATATTTGCAA	AAATTCAACI	GTTTTATCAG	GCTATCTTT
226921	TAATAATATA	TTCATTTTAT	CTATATCTGA	GGTTTTAGCT	TCTTTGTACT	TCTGACCCAA
226981	TTGCATGTGT	GCTTTCTTTC	TCCTTCATTA	GACTACTTAG	TCATTTACTA	ATTTTAAGAA
227041	TAGCTTGTCT	TTTATTTATT	TACTTATTTA	TTTTTGAGAC	GGAGTCTCAC	TCTGTCACCC
227101	AGGCTGGAGT	GCAGTGGCGC	GATCTCGGCT	CACTGCAACC	TCCGCCTCCC	GGGTTCAAGT
227161	GATTCTCCTG	CCTCAGACTC	CCGAGTAGCT	GGGATTACAG	TCATGCACCA	CCATGTCTGG
227221	CTAATTTCTG	TATTTTTAAT	' AGAGATGGGG	TTTTGCCATG	TTGGCCANGC	TGGTCTCAAA
227281	CTCCTGACCT	TAGATGATCT	ACCCACCTTG	GCCTCCCAAA	GTGCTGGGAT	TACAGGCATG
227341	AGCCACTGCG	CCCAGCCCTG	CTTGTCTTTT	TATTTTATAT	TTGATTAGCT	TTATCTTTTA
227401	TCAAGCTTAT	GTCCTATTTC	CCTTTGCTTT	ACTTCATATA	V Duning Transfer	TGGATAGTTT
227461	ATTTATTTT	CATTTAATTA	TGAAACAGGT	TAAAGCTTAG	AGGAAAATTG	CTCCTCTAAG
227521	TCCACTTTTG	TGGGCAGATT	ACATTTTGCT	GTGTTGTGCT	CCCAAATTCA	TIGITCITIT
227581	AATGCTTTAT	TTCTCAAGTT	AATAACCTAT	ATAGTAAAAA	AGTGGCTGTT	GACTCTCAGC
227641	TTTTTTTTT	TTTTTTTTT	TTTTTTTGTA	GATACAGGGA	TOTTGOTGIT	TTGCTCAGGC
227701	TGGTCTGAAA	CTCCTGGCTT	CAAGGGATCC	TCCTGCCTTG	GTCTCACAAA	ATGCTCAGGC
227761	GACAGACATG	AGACACCATG	CCCAGCCATG	TCTCTCTCCT	TATATATATA	AAGAAAACAG
227821	ACACACTGAG	GCATCCTATC	ATCTCACTCT	TGGTTTCACT	PUTCTTCTCT	GGAAGTTTTG
227881	CTCTGACCTT	TTGCAGTTAA	TGTATTAATT	TTGCATTGAG	TAGTTTCCAT	AGAAGAATTA
227941	TAGCATTTGC	ATTCTGTTGG	GTATTATACT	TTTCACTGTT	ATTTGDACAT	AATTTGAGGG
228001	CTGAAACCAA	GATGAGGCAA	GTGAGGTGCC	CAGGAAGCAA	TATTTAAGGA	AATTIGAGGG
228061	CTTAGGCTCA	TGCAAGAACA	GAATTGGCAC	ATGAGAGTGA	GTGCCTCCTT	AARTECTT
228121	GCTGGACACT	TCTTGCTCAC	TTAGCATACC	CCTCCACAAT	GAAGTGTTTT	AATTTTGAGT
228181	TTTTCATGTC	CATCCTTTAT	CCTTCTTCAT	CTCDDALCAT	TTCAATGGAG	TIGITITGIT
228241	GAGCAGTACT	TGGATGAGCC	TCTGAGTCCC	ACAGTAGCTG	AGAATTTATT	TORTHOR
228301	CTTTATGATC	ACTGTGGAGC	CTTAAAACAT	TGTAATATTA	ACTTAGCTGG	CARGACA
228361	TTTGTTCCAC	AATTTGTCTT	ATTCAGAACA	GTATTGACTT	CCTGCTAGTC	TCTTCTTCTCA TC
228421	TCCAATATGA	GGAAGTCTAG	TTAGCCAGCT	ACTITUTETA	GGAGAGCTAT	CTTTTTTCTGATG
228481	GGTGCTATAG	GATTCTCTTT	ATCCTGGAAT	TCCTTCACCA	AGATGTGCCA	ACCTCTTA A
228541	CATTTTCTCT	TGCTTTTTGG	CTGGTGGTCT	TAGAGTTTCC	TTCGATTTTG	TTTTTT TTTTT
228601	TGATTGTCCT	CAATTTGTTT	TCTTTACTAA	GAATCTCTCT	TCTATTTATC	TOTALLIA
228661	AACCTTGTTG	CCCATCTTTC	TGGTTTCTGC	TGACTTTCAT	TTTTGGACCT	TTTTACTTTCC
228721	TTTCTCCATG	GACTTTTTGG	TAGTGGAGGC	AGGCAAACAC	TTTCCAAAGT	CETTOTT
228781	TTCCATCAAT	TTCAACTTAT	TTCCTAAAAT	TGCCTCAGAA	TGTGCCTATG	TCCACAAT
228841	CCCTCCTTCC	ACTTTAGAAA	GGAAAGGCAT	CCACACTTTA	TTTAGGTGCA	ATTCCACAATAT
228901	TGTAAACACT	TTCTGGTTGT	CAACAAAGGA	GTACTTCCAA	ATATTGGTTT	GGGGATA
228961	TGCTAATGAT	TAACACATTC	ACCTTGGCTC	TTGGTTTGCC	TGCTCCCTCT	TCTTTTT TCT
229021	GCTGTGTGTA	TTTTTTTAA	TCACTGAGAA	TATGCACAGT	ATTGTATGTT	TCITTATCT
229081	AGAGGACTGG	CCAGAGTGGG	AATGTTCTGA	ATTCAGAATA	ACTGARGORG	TIMITALAMG
229141	GAACTCATTC	TTTCAAATGA	AGCTGGCATA	TTTTCCCAGA	CCACCAAACAG	TACAGGAIAG
229201	TTTAAAAAAC	TTGATATGAA	TGATACAATA	AAGTGGTTAG	AACTTTTATT	TCWWINIWIA
229261	TATGTCATGA	AATACTTATT	CTAATTATAG	TCACTCTTCA	TOTTE	TCTTRTRACT
229321	TGTTTAATGT	TITCTTTTAT	TTACAAAACA	ATTTATTT	TCATCAAAAC	TCIIMIMMCM
229381	CAAGTTAAAA	ATATTCAAAG	GAATGCCTAA	AGTTTTCAAA		TITINGMAMI
229441	ATCAAAAGAG	TCTGAAGACC	ATTTAGCTAT	CCAAATTGTT	TATTTTTA	CAGTATCCCT
229501	TCTAATATTT	ACTATTTATA	ATCCTTAAAA	ATTTGCCTTA	GCACAGGAGA	ATTGCTTCX N
229561	CCCAGGAGAC	GGAGGTTGCA	GTGAGCCAAC	ACAGTGCCAC	TGCCCTCCAG	CCTCGGCGAC
229621	AGAGTGAGAC	TCTGTCTCAA	AAAAAAAAA	AAAAAAAAA	AAAAAAGGCC	TRACOCARG
229681	AAACAAACAA .	AAAAATCCGC	CTTAACATTA	TTTGTTCATT	AAAAACTTTC	מידי ביד ביד בידידי
229741	CTAGTTTCCC	TTTCCTCTCA	GCCCATTGTC	ATATTTTGAT	TTTTATCACT	recomposite and the second sec
229801	GACATATGAG	GTTTTTGTTT	TTTTTTTTT	TTGGAGATGC	AGTCTCCCTC '	TGTTGCCCGT
229861	GCTGGAGTGC	AATGGCGCAA	TCTTGGCTCA	CTGCAACCTC	TGCCTCCTGG	מתרבא בכרב א
229921	TTCTCCTGCC	TCAGCCTTCC	AAGTAGCTGG	GATTACAGGC	ACCCACTACC	ACGCCTGGCT
229981	AATTTTTGTA	TTTCTGGTAG	AGACGGGGTT	TCACCATGTT	GGCCAGGCTG	TOTOGRADOR
			<b></b>			

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230041	CCTGACCTC	A AGTGATCCA	C AATCCTTGG	CTCCCAAAG	T GCTATCATT	A CAAGCATGAG
230101	CCACCTGCC	C AGCCAGAAT	A TATGTTCAT	TTGAGTCCT	TABOARACTO	C ATAAGAATTT
230161	TAGGAATTC	A GTTACTTTC	T TGAGAAAAT	TOTORDICE	TCCCAAMAGI	TTGTAGCCAA
230221	TTATATTGA	T TTCTCTTTT	T CATATTGAG	ייייייייייייייייייייייייייייייייייייי	T AAAAACTEE	TATGTGTGAA
230281	GATTTTTGC	A CTGTAGTTA	A AGAAACCAC	TGTGTGTTG	TTDDCCCDm	A AGTACATGTA
230341	TTCAAATAA	A TTGAGGTGG	GTTACTCTG	GAATCAAAG	2 DADACCECAI	A GAAACAGGCA
230401	GCCTCAAAA	G GTCTTAGCT	TAGCAACTTO	CTCCATTGT	T CARACCIGAL	GAAACAGGCA GGCTTGAACT
230461	TGTATTTTC	C CTCTACTCAL	A CATTTAAGG	CTCAGAAGA	L VVINUTVINUT	GGCTTGAACT GTGAAATTTA
230521	AGTAAAGTG	C TCACTCTTT	C GCTTTAACA	ACCCTAGAGE	COTCOTACO	GIGAAATITA AGAGCCTCAA
230581	CAGACCGTT	T TAGCTTCCA	AGGGAGTTC	GGACACCATO	ATTCACAROGC	ACAATACATC
230641	ACACATAAT	T GAGAAAAGAT	AGTTCCACCA	AATAAAGTTO	ALICACOMON	ACAATACATC
230701	AAGAAATCT	T GGAAATAGGT	מממדמדמדד ז		CCTTTTTTTT	TGTTATGGAA
230761	TAGGACCAG'	TCTACTTAAC	CCACCCATTT	CCCDDDDTD	ACTCACARMO	GTTTCTTTTG
230821	GGGACTCCT	TTTGTAGCT	CAAGTGCCAC	TABCARTECT	TAGGAGAATC	GTTTCTTTTG
230881	AGGTGATTT	AGTTAATAT	ATCAATTATT	י דרמערטינון י	CCCTCTART	TGCAGAGGGA
230941	ACGGAGCCC	TCAGCATTCC	CTGCAGGGAA	CTCCACTCC	TOUCICIAAIG	TGCAGAGGGA
231001	AGCTTTCAA	TGTTTTGAA	TCACTTTCAC	GETECTENTO	TITIMICAAC	TTGAACAGCT TTTGAAATCA
231061	GAAGATGATT	CTGCCTCTT	TAATATGTGA	CTCCTCAGAT	TAGIIGCTTT	CTCGCTAGTC
231121	TTAAGAGTG	ATTACCCTC	GTGGTCCAGC	CICCICAGAI	CCAGAAAGIG	CCCTATCCC
231181	TGGGGGAACT	TATCAGAGAAA	TTGGTGCCAT	GCITAIGAAC	CCACATCTAA	GTGAAGCAGA
231241	GAGCCCCGC	TGATGAAAAT	CAGTGGACAG	CAUCATAACA	GGAAGGCACA	GTGAAGCAGA TAATCACCCA
231301	GGAGCATGA	AATCCAGGCC	CACIOCACAC 'ARTCTCCCAC	CATCALIALL	TACAACTITG	GGAGTTCTTG
231361	GAACCGATTC	TGATGAATGA	CTGTTTAGCC	ATTITION	AATTITTGTT	TGGCTGCTTG
231421	CATACAGAGG	TTGGATGTAA	ACGGGCCTTT	CCCCTCTCTCTT	GIGGCATACG	ACAGGAACTA
231481	AACTGTGTCA	CATAGGTTCC	AAATGGTGGC	CTCAATACTA	TTTT CD D COD	ACAGGAACTA AGGTACAATG
231541	AAATTGAGTA	AGTCTTTTCC	TCTTTTCCAC	ATACCATCAT	TATTCAMCIA	TTTCTTCAAA
231601	GTTAACTATT	TGTATTTGGT	ייע מיידידידע איי	ACADATCTAN	TATICATATA	TITCTTCAAA
231661	TCTTTAGTCT	TAAGGTTGAT	GCTCTCCATG	TCCTTCCAAA	AAAAACCTATC	TCAAGTTTAG
231721	TATATCCTCG	CCTTCAGATG	GGATTATTCC	TALLALCAN	TTCTTT ATAT	TIGCTITIAT
231781	CCACTTTTTT	TGTGGCTCTG	GGTGAGATGC	TATACCTACA	ATCACA ACTIC	ATACTTTGAG
231841	TGTCCCTGTC	ACAAAAGTGG	ATAGCCTAAG	TGGTGACTTT	TACCTCCACT	ATACGTGTGT
231901	GTATCACACA	CCAGCCGTAT	GCCAGGCACC	ACTCTAGGTG	CTACCCCACI	CCAAATATAT
231961	AGACAAATGC	AACCCCTGCC	CATGTGAAAG	AGAATAAGAC	DATABAGRIAC	AGCAGTAAAC
232021	GTTATATGGA	GGTGGCAAAT	GCTAAAAAGA	AAAATTAAGC	DECEDIANCE	ACTORMOGRA
232081	AAGATGACAT	TTGGGTAAAA	GCCCATGTAT	ATATGTTCTA	TTCCTTTT	TTCTCTCCAA
232141	AGCCCTGACT	AATACACAAT	GACTITGAGA	AGTTACTGGC	מינייניים מינייניים אינייניים אינייניים אינייניים אינייניים אינייניים אינייניים אינייניים אינייניים אינייניים	TCACACTATE
232201	CGGAGTGCTG	AGAGCCTTCT	TAGTGTGTAT	TCAGTGTTTT	AAGAGAGCTT	CTCCATCAAT
232261	AATAAATAGG	ACAAAATTTA	TCCAAACTTA	AGCCTTGCTT	TAGGTAAAAG	CCCTCCTCTT
232321	ACAAGGTAGA	AGGTTATTAT	TTGACATTTA	AATCCAACTG	AACACTAATA	ACACTA ATTON
232381	ATTAAAAGTT	TTTAAATCAC	AACTGCGTGC	AAAATAAATG	GAACTGCCAT	CCTCCCCAAC
232441	IGIGCATGAG	TGGTGTGCAT	GGGAGACAGC	ACGAAGCTAA	TCCCACTCAT	CTTCC A CCTT
232501	GCTCCATTTT	TCTCCTAAAA	TCAGTAAGAC	AGAAGCTGGT	CAGATTATCA	DEDECECTAC
232561	TIMESCACAG	CAGTAGCATT	TGGAAGGGGT	TGCTCTCATT	AGGCAGTGCC	TCACCACAAC
232621	AAGAGATGAA	CAAGCCCTGT	ATCTGAAGCC	ATCATGCCTA	GTTATGGTCC	CCGACCACAAC
232681	AIGAIGCCIG	GAAGGGAGGC	CCCCTGCACC	CTAGAAAGCT	GGGTGGGTTC	TACTCTCTCC
232741	TTTACTGCTA	AAAACCCTCT	TCTTTGGATC	TGGACTTTAC	CTCTATCTCA	TACIGICIGC
232801	TAATATATGA	TTTGGCACTG	AGTCTGTCAC	TGCTGCTAAC	TCAGCAGTTC	TACCOTOSTO
232861	GCCCCATTGC	CTCACAGAAA	GAATTTCATA	GCTTCCAGCA	TCCTCTCTCC	ጥጥሮአጥጥአጥአሮ
232921	TITGATTTCA	GCATTGCTAT	TTTTTCTCTT	GGGTGTTGCA	GCTCTCTCTC	TO COMPONE TO THE
232981	GICTIGTIGG	TTTTCTGCTA	ACTCCTGCTT	TTTTTCTTTT	تالملسلس بالملسلسل	AGACGGAGTC
233041	regriciere	ACCCAGGCTG	GAGTGCAGTG	GCACAATCTC	GGCTCACTGC	AACCTCCCCC
233101	TCCCGGGTTC	AAGCTATTCT	CCTGCCTCAG	CCTCCCAAGT	AGCTGGGACT	ACAGGGGGGTC
233161	ACCACTATGC	CCCACTAATT	TTTGTATTTT	TAGTATTGCT	GTCATCAATC	CACATGTCCA
233221	GAAGCACCTA	GAAACTCTAA	TTCTTTGTAG	GTATCAAACC	CTAGGACTCT	TTCATGICCA
						LICCICIAMI

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233281	CACAATATAT	AATCCCTGAT	TCCCAAACAC	GGTCTTTCA	TATACATTT	r ccactgtaca
233341	TACTTTCTGA	L CCTGGAAAGC	TCTTACACAZ	ACACGCCCTC	CCCTAGGAAC	מממידמידים ב
233401	TGTTCCCAGG	AAGAATCAGI	' CACCCAACAG	TGTCCTTGTC	ACATOTTAGO	ב ייייריזיאראריריי
233461	TTATTTGTTC	TATCTGAATG	TAATCTCCCA	GAGGGTGTTA	TCATCTTTT	ר יייייייייה אמאיים
233521	GAGTCTTGCT	TTGCTGCCCA	GGCTGGAGTG	CAGTGGCATG	ATCTCGGCTC	CACAGCAACCT
233581	CCACCTCCTG	GGTTCAAGTG	ATTCTCCTGC	CTCAGCCTCC	TGAGTAGCTO	GGATTACAGA
233641	CGTGTGTCAC	CACACCTGGC	TAATTTTTGT	ATTTTTAGTA	GAGACAGGG	TTCACCGTGT
233701	TGGCAAGGCT	TTCCTCGAAC	TCCCAAACTC	AGGTGATCCA	CCCACCTCAC	CCTCCCAAAG
233761	TGCTGGGATT	ACAGGTGTGA	GCCACCATGT	CCAGCCCCAT		TAGTTTAGTT
233821	CTTAACAAAT	AGTCTGACAC	AAAGTGGATA	TAACAATATT	יייי איייי איייי איייי איייי איייי איייי אייייי אייייי אייייי אייייי אייייי	AATAACTAAA
233881	TGAATATTTC	CAGATTTCCT	GGTGCTCTCA	AAGTTTTATG	TTOCARITATO	AAAACAAGTC
233941	TAAAATACCT	GCCTCAAGTT	TTTATCTGTA	CTATGATTTC	TINCHANAGE	AAAAACAAGTC
234001	GGGGTAAAAA	CTGAAACAGG	AAATACATAT	AACTGAAAA	TTTTTCCHAALA	TTAGTATGAT
234061	AATACTAGGT	CATTTTTCCT	GTTTCCCCAA	CTTCATTTC	TATECTATE	AAAAGAAACA
234121	AGTAAATGTA	TGTTAATTTA	ATTTAAAAGA	AGTAGTCTAC		GTTAAAAAGA
234181	AAAAAGTATT	TTAAAAAATT	ATCTCTGGAA	GGATACACAG	GGAACATTCC	TCTGGTTTCT
234241	TCCAAGAGAG	AAATGAGGAA	CTAGAGAGCA	TGGCCAACTG	GCGTTTTTCCT	TTTGTTTTTG
234301	TTTGTCTATC	TGTTAGCTTT	TTATTATT	CTTTTCTAGG	TTTC A ATTTC	AAACCACATA
234361	AATCTGTTAC	ATGCTCATAA	TAATAAGTTT	CITITOIAGG	TTTTTCCCCCC	GTGCAATGAC
234421	TTACACCTGT	AATCCCAGCG	CTTTGGGAAG	CACACCTCCC	1111GGCTGG	AGGCCAGGAA
234481	TTTGAGATCA	GCCTGGGCAA	CATAGTGAGA	CCCTCCCTCT	AGGATACTTG	ACAAAAATTA
234541	GCTGGATATG	GTGGTGCATG	CTTGTACTCC	TAGCTACTTC	GIAGAAATAA	GCAGGAGGAT
234601	CCTTTGAGTC	CAGGAGTTTG	AGGCTGCAGT	GAGCTACIIG	CACCCACTICAG	ACTATAGCAT
234661	GGGCAATAAG	GTGAGAACTT	GTCTCAAAAA	AAAAAGGGGG	CCCCCACIGC	AATAAATAAA
234721	TATAAACAAA	ACTTTTGTTT	CAAAATATGT	AATATTTACC	ACTENANCA	TCTGAATTGT
234781	AGAGCTAAAA	AGTACTTAAA	AGTTAATAAC	TATTCTCTCC	TOTALAGRA	TTGTTATCAA
234841	AGTATAATTT	TTATCCAGAA	AATCATCCAT	ATCAGCAAGC	TITARAGAA	CAAAATGACA
234901	TATCCATGTA	ATTAGCTCCC	AGGTAATTAG	CAGGCAGCCT	CTACTCACCT	TGAGTATTCC
234961	TAATCTAAAA	ATTGGAAATT	CAAAATGCTC	CABANTOTO	A A COMMONDON	ATGCTAACAT
235021	GATTCTCAAA	GGAGTGCTCA	TGGAGTATTT	CAGATTTTGG	AMCITITION	AIGCTAACAT
235081	CAGTATAATG	CAAACATTCC	AAATCTGAAA	AAATCTCAAA	TACTTCTCC	TCTAAGCATA
235141	AGGGATACTC	AACGTGTGTT	AGCTAATTAG	ACCOTTCATC	CTCTCTTCTT	CACCECATA
235201	TCTTCAAGGT	AACCTCTATC	CTCACTTCTA	ATACCATCAA	COTTOTICIA	GACCTCAGCT
235261	TTGGATTTTC	AGGAAAGTTG	CAAAGATAGT	ACADAGACAC	TACACCACAC	TTAGAATAAT
235321	TCTTTCACCT	AGCTTTCCCC	CATTGTTAGG	שריישריישריישריי	ATTATCATAC	ATTOCCATATA
235381	TATAAGCAAC	TCACATTGAT	ACATGAAACT	CTATTARCELL	ATTAIGATAC	ATTIGICAAA
235441	TTTCACCACT	GTTTCCACTA	VACALALIA LA	TCTCTTCCAA	CCTCC A A TCT	TTTATGTGGA
235501	ACTGCATTTT	CTTGTCATAT	CTCCCTAGTC	TOTALICOM	TOTOLOR	GGAATACCAC
235561	TTCTTGCTTT	TCATGACCTT	AACAGTCCTG	AMBATCATTT	CCTTTTTTTTTT	TCTCAGTCTT
235621	CCGGAGTTAT	AGATTTTTTG	AAATAATACC	ACAAGGGCAA	AGGGCCCTTC	TTCTCT CATC
235681	ATTTTAGGGA	GAACATGATA	TCCACATGAC	ATCACTCATA	TOUGGCCCIIC	TOTCACATC
235741	TAGGTAATGT	TTCAGGTTTC	TCTACTGCAA	AGTGATTTT	TIMECTICA	TERCOCCRO
235801	TGAACTTATC	AATTTTGTTT	TCTTCCATGA	CTAATACTT	TICCCIIAMI	CCTARARA
235861	CATTGGGGCC	AAATCTTAGA	TCATGTAAAT	THEFT	TGTIVIIVIV	A A A COURGE
235921	AATGTTTGAT	ACATTCTAAA	AGATGTAATG	TTTGATACAT	TACATCTACT	AAAAGCTTGT
235981	ATTTTTAGTT	ACTITTGTAT	AAGGTGTGAG	AGATGTCTCC	TWOWICINGS	CCTTTGATTT
236041	TTGTGGTGTT	CCAGTACTAT	TTGTTGCTAA	GACTATOTO	TTTCACII	TATTAACACA
236101	CTTAGTTGGC	AATATTTTTG	TTGGTTTATT	TCTAGACTGT	TTCCALIGA.	COLORCAMO
236161	GTGTCTATCT	TTTTGACAAA	ACTGTTGATT	ACAGTANGOT	TIMICICALT	TCAUTGATTT
236221	TGTCAACTTG	ACTGAGTCAG	GGGATAACCA	GCTATCTGGT	TAAACATAGI	TTCTCCCCCCC
236281	GTTTGTGAGC	GTGTTTCTGG	ATGAGATTAG	CCTTTGAATA	GGTGATCCTA	CTANACTANA
236341	CTGTCTTTCC	CAGTGTGGAT	GCATTATGC	CACCTGATAT	TCAGGGTCTG	DATACARCIANA
236401	AMGGCAGAGG	AAGGGGGAAT '	ITGGGCCTTT '	TTTTCTGCCT (	CACTGCTTGA	GCTCCC A CAT
236461	CTCATCTGGT	CTCCTGCTCT	TGAACTGGGA	TTTACATCAT	CAGTTCCTCT	GCTGGGACAT
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236521	CCTTCAGATT	CAGACTGAAT	CATACCACCA	GCTTTCCTGG	GTCTCCAGCT	TGCAGATTAC
236581	AGATCATGGG	ACTCCTCATC	TTCCATAAAT	GCATGAGCCA	ATTCAGTCTA	TGTCCTTGAA
236641	AACTGCCCCA	CTGCAGATTA	AGGCTTTTTT	CCACTAGGTG	AAATAAAGAA	GCTTGTTAGA
236701					CAACACATTG	
236761	AGTGCAGGGG	TGGGGATGAG	GGTATAGTCC	TCTTGTTTGC	TGAGAAGAGA	ACTGTATTGG
236821	GAAAGCTCTA	GAAGTGTTTG	ATACATACAT	AAACAAGGCA	TGGTTTTTGC	ACTTAATTTC
236881	ACATTACATT	TTTCCCAGAA	AAAAAGGAAT	GTATAGGCAT	CACGTAACTG	TACTAGCTGG
236941	AGTCATTCTT	CCTGATTATC	AAAGGTAAAC	AGTTATTAAT	CCTATACCAA	GATGTCAAGG
237001	AGAAGTACTT	TTGGAACACA	AGGAATTCTC	TGGGAGTCCT	TACTACTCTC	AAGCCCAGTG
237061	AAAAAGTTAA	TGAAAAACTA	TAGTACCTTC	CTATAAGCTG	GATGACTAAT	TACCAGGCTC
237121					TGAGGTGCTG	
237181	AATGGAGCAT	AGAATAAGAG	TAGTAAAGAA	TGCCAAAAAT	GCTGTCATGT	ATCCATTGAC
237241					CTGTTACGTA	
237301		TGTGTGTGTG				

Figure 9 (Pag 74 of 74)

SUBSTITUTE SHEET (RULE 26)

International application No. PCT/US97/17658

IPC(6) US CL	ASSIFICATION OF SUBJECT MATTER : C07H 21/04; C12Q 1/68; C12N 15/63, 15/85; : 536/23.5; 435/6, 70.1, 325, 320.1			
	to International Patent Classification (IPC) or to be LDS SEARCHED	oth national classification and IPC		
<del></del>	documentation searched (classification system follo	wed by classification symbols)		
U.S. :	536/23.5; 435/6, 70.1, 325, 320.1	wa by classification symbols;		
Documenta	tion searched other than minimum documentation to	the extent that such documents are included	d in the fields searched	
APS, DIA	data base consulted during the international search ALOG'S BIOTECH cluster. ornatosis, BTF1, BTF2, BTF3, BTF4, NTP-3, NT			
C. DOC	UMENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to claim No.	
A, P	RUDDY, D.A. et al. A 1.1-Mb tra hemochromatosis locus. Genome Res 5, pages 441-456, see entire document	1-20, 22-77		
X	FISCHER, L. et al. Cloning of the basic transcription factor BTF2. Scie 257, pages 1392-1395, see entire doc	28-33, 71		
X	MARGOTTIN, F. et al. Participal transcription of the yeast U6 gene by 25 January 1991, Vol. 251, pages 42.	RNA polymerase C. Science.	22-27, 70	
X Furth	er documents are listed in the continuation of Box	C. See patent family annex.	;	
Special categories of cited documents     document defining the general state of the art which is not considered to be of particular relevance.		"T" later document published after the inte- date and not in conflict with the appli- the principle or theory underlying the	cation but cited to understand	
E earlier document published on or after the international filing date document which may throw doubts on priority claim(s) or which is cited to establish the publication data of another citation or other		"X" document of particular relevance, the considered novel or cannot be consider when the document is taken alone	claimed invention cannot be ed to involve an inventive step	
special reason (as specified)  document referring to an oral disclosure, use, exhibition or other means		"Y" document of particular relevance; the considered to involve an inventive combined with one or more other such	step when the document is documents, such combination	
P* docu	ament published prior to the international filing date but later than priority date claimed	being obvious to a person skilled in the "&" document member of the same patent		
Date of the a	ctual completion of the international search	Date of mailing of the international sea	rch report	
20 JANUARY 1998		1 2 FEB 1998		
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231		F. PIERRE VANDERVEGT		
acsimile No		Telephone No. (703) 308-0196		
rm PCT/IS	A/210 (second sheet)(July 1992)*		\ / /	

International application No. PCT/US97/17658

C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT			7658	
Category*				
	Citation of document, with indication, where appropriate, of the relev	Relevant to claim N		
X	ZHENG, X.M. et al. Sequencing and expression of con DNA for the general transcription factor BTF3. Nature 1990, Vol. 344, pages 556-559, see entire document.	34-39, 72		
X	PANTEGHINI, M. Electrophoretic fractionation of 5'-n Clinical Chemistry. February 1994, Vol. 40, No. 2, page see entire document.	52-57, 75		
	BURT, M. J. et al. A 4.5-megabase YAC Contig and ph map over the hemochromatosis gene region. Genomics. I 1996, Vol. 33, No. 2, pages 152, 169	1-6		
	1996, Vol. 33, No. 2, pages 153-158, see entire documen	ıt.	7-20, 22-77	
1	VERNET, C. et al. Evolutionary study of multigenic fan mapping close to the human MHC Class I region. J. Mo November 1993, Vol. 37, No. 6, pages 600-612, see abstr particular.	1-20, 22-77		
			-#-	
DCT/ICA ICA	0 (continuation of second sheet)(July 1992)	1		

Form PCT/ISA/210 (continuation of second sheet)(July 1992)\*

International application No. PCT/US97/17658

Box ! Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)	
This international report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:	
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:	
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:	
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).	
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)	
This International Searching Authority found multiple inventions in this international application, as follows:	
Please See Extra Sheet.	
1. X As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.	٥
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.	
As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:	3
•	
No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:	
The additional search fees were accompanied by the applicant's protest.	
No protest accompanied the payment of additional search fees.	1

Form PCT/ISA/210 (continuation of first sheet(1))(July 1992)\*



International application No. PCT/US97/17658

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

This application contains the following inventions or groups of inventions which are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for all inventions to be searched, the appropriate additional search fees must be paid.

Group 1, claim(s)1-20, drawn to polynucleotide sequences containing at least one polymorphic site, polypeptides encoded thereby, antibodies to said polypeptides and a method to determine the presence of the HFE gene mutation.

Group II, claim 21, drawn to the lymphoblastoid line atcc crl-12371.

Group III, claim(s) 22-27 and 70, drawn to BTF1 nucleic acids, gene products, vectors and antibodies.

Group IV, claim(s)28-33 and 71, drawn to BTF2 nucleic acids, gene products, vectors and antibodies.

Group V, claim(s) 34-39 and 72, drawn to BTF3 nucleic acids, gene products, vectors and antibodies.

Group VI, claim(s) 40-45 and 73, drawn to BTP4 nucleic acids, gene products, vectors and antibodies.

Group VII, claim(s) 46-51 and 74, drawn to BTF5 nucleic acids, gene products, vectors and antibodies.

Group VIII, claim(s) 52-57 and 75, drawn to NPT3 nucleic acids, gene products, vectors and antibodies.

Group IX, claim(s) 58-63 and 76, drawn to NPT4 nucleic acids, gene products, vectors and antibodies.

Group X, claim(s) 64-69 and 77, drawn to RoRet nucleic acids, gene products, vectors and antibodies.

The inventions listed as Groups I-X do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, they lack the same or corresponding special technical features for the following reasons: Groups I and III-X are drawn to physically different genes and their gene products and each therefore constitutes a separate invention. The lymphoblastoid cell line of Group II is not dependent upon the vectors of any of the Groups I and III-X and therefore constitutes a separate invention. Accordingly, the claims are not so linked by a special technical feature within the meaning of PCT Rule 13.2 so as to form a single inventive concept.

Form PCT/ISA/210 (extra sheet)(July 1992)+